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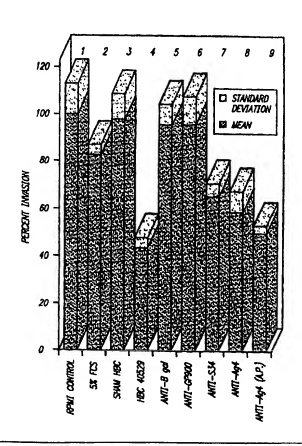
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(54) Title: VACCINES, ANTIBODIES, PROTEINS, GLYCOPROTEINS, DNAS AND RNAS FOR PROPHYLAXIS AND TREAT-MENT OF CRYPTOSPORIDIUM PARVUM INFECTIONS

(57) Abstract

Vaccines, antibodies, proteins, glycoproteins, DNAs and RNAs useful for passive or active prophylaxis and treatment of *Cryptosporidium* infections. *Cryptosporidium* antigen comprised of a protein with or without carbohydrates attached thereto. Polyclonal and monoclonal antibodies directed against the antigen. DNA and RNA encoding the *Cryptosporidium* antigen, mutants, variants and fragments thereof.



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VACCINES, ANTIBODIES, PROTEINS, GLYCOPROTEINS, DNAS AND RNAS FOR PROPHYLAXIS AND TREATMENT OF Cryptosporidium parvum INFECTIONS

This invention was developed partially with U.S. Government support under National Institutes of Health Grant Nos. AI-29882 and AI 30295. The U.S. Government may have certain rights in this invention.

This application is a continuation-in-part of the US Application Serial No. 08/415,751 filed on April 3, 1995 which is a continuation of Ser. No. 08/071,880 filed on June 1, 1993 which is a continuation-in-part of Ser. No.07/891,301 filed May 29, 1992.

BACKGROUND OF THE INVENTION

Field of the Invention

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This invention concerns vaccines. antibodies. proteins, glycoproteins, DNAs and RNAs for prophylaxis and treatment of Cryptosporidium or Cryptosporidium infections. In particular, this invention concerns a Cryptosporidium antigen comprised of protein with a carbohydrates attached thereto, as well as polyclonal and 25 monoclonal antibodies directed against the Additionally, the invention concerns DNA and RNA encoding antigen, mutants, Cryptosporidium variants and fragments thereof.

Background and Related Disclosures

The genus Cryptosporidium consists of Apicomplexan parasites that invade and develop within epithelial cells of the gastrointestinal, hepatobiliary and respiratory tracts of a wide variety of vertebrates including reptiles, birds and mammals. Cryptosporidium was recognized as a cause of animal disease for several decades before the first cases of human cryptosporidiosis were reported in 1976. However, it was not until 1982 that the magnitude of the disease caused by this parasite in both AIDS patients

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and immunocompetent hosts began to be appreciated. Subsequently, Cryptosporidium has been found to be among the top four causes of human diarrhea worldwide, and to be an increasingly recognized cause of diarrhea in children, animal care workers, and travelers. (Cryptosporidium in Humans, Ed. J.P. Dubai et al., CRC Press, Boca Raton (1990)).

Large waterborne outbreaks of cryptosporidiosis caused by contaminated municipal water supplies in the US and in the UK have been noted in the last ten years (N. Engl. J. Med., 320:1372 (1989), and 33:161 (1994)). The most recent outbreak in Milwaukee in April 1993 involved 400,000 persons and led to the subsequent deaths of more than 100 immunocompromised persons. Like a number of other waterborne outbreaks, the Milwaukee outbreak appears to have been due to contamination from farm or abattoir runoff and specifically due to cryptosporidiosis among cows/calves. Nosocomial transmission in hospitals from patients to staff, patient to patient, and contaminated ice to patients and staff have also been well documented (J. Inf. Dis., 158:647 (1985)).

Waterborne and nosocomial spread reveal a number of biological characteristics of oocysts. First, infectious dose of the parasite is very low. The ID, for human volunteers with normal immune systems is 132 oocysts (New Engl. J. Med., 332:855 (1995)). Second, infected hosts, for example calves, excrete large numbers of oocysts; on the order of 1010/day. Third, the oocysts are fully sporulated and ready to infect when excreted. Fourth, the oocysts are environmentally hardy; they remain infectious in cool, moist areas for 3-4 months, and they are not killed by chlorine levels permissible in drinking water. Fifth, the oocysts are quite small, 4-6 μ m, and are thus difficult to filter.

The clinical importance of cryptosporidiosis has increased markedly with the recognition of a lifethreatening form of the disease in patients with immunodeficiency disorders such as AIDS, hypogammaglobulinemia, and chemotherapeutic

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immunosuppression. The prevalence of cryptosporidiosis in AIDS patients in the US is estimated to be 5-10% and in central Africa 40%. Immunodeficient patients may have fulminant cryptosporidial diarrhea that may persist until death, whereas the diarrhea of immunocompetent patients is self-limited and rarely lasts more than 2-4 weeks. Cholera-like diarrhea is common in immunocompromised patients with reported losses of up to 17 liters of water Hepatobiliary disease may result in severe per day. abdominal pain and nausea. Removal of immunosuppression in chemotherapy patients leads to resolution of the diarrhea. Occasionally, AIDS patients with cryptosporidiosis will be able to eliminate the parasite after initiation of antiretroviral therapy (Am. Intern. Med., 116:840 (1992)).

Among those who develop the disease, a quarter have CD4 counts greater than 209, suggesting that the cryptosporidiosis disease may appear relatively early in the course of HIV disease. Unfortunately, few details about the biology of the organisms and the molecular mediators of the disease process have been described and no effective therapy has been discovered.

infective forms of Cryptosporidium, sporozoites and merozoites, adhere to the host cell and release the contents of anterior organelles (rhoptries, micronemes or dense granules) during the invasion process (Parasitol. Today, 8:28(1992)). Proteins involved in these events have in many instances been found to be the target of invasion blocking immunity in vitro and neutralization in vivo (ibid). Active and passive immunization studies using malaria and Toxoplasma challenged or infected hosts, have shown that certain secreted components of the apical complex organelles are the target of protective antibodies in these related Apicomplexan parasites. In some cases, as for example in the case of the circumsporozoite and merozoite surface proteins of malaria, these antigens are under development as vaccines.

While the actual interaction between Cryptosporidium and the host's immune system is poorly understood, it is known that disruption of either the cellular or the humoral

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components can result in protracted cryptosporidiosis (Parasitol. Today, 8:24 (1992)). However, specific antibodies alone neutralize the organism's infectivity. In vitro and in vivo observations indicate that antibodies to Cryptosporidium parvum inhibit invasion and intracellular development leading to protection in challenge experiments, or amelioration of infection in established disease (Infect. Immun., 59:1172 (1991)).

One source of such antibodies is hyperimmune bovine 10 colostrum (HBC) collected from COWS immunized Cryptosporidium oocysts. Calves challenged with Cryptosporidium oocysts are protected from the development of the disease by the administration of HBC (Infect. Immun., 61:4079 (1993)). Some immunocompromised AIDS patients infected with Cryptosporidium have also responded 15 to HBC with a reduction in, or disappearance of, the symptoms of the disease (Gastroenterology, 98:486 (1990)). Immunoglobulin from HBC (HBC Ig) has been found to inhibit the ability of the sporozoite to invade and/or develop 20 intracellularly in vitro and it has been used immunoprecipitate at least 22 different surface radioiodinated proteins of Cryptosporidium sporozoites. Western blot analysis of proteins of whole oocysts, which contain sporozoites, indicates that HBC predominantly recognizes two proteins of sizes 250 kDa and >900 kDa 25 (Infect. Immun., 61:4079 (1993)).

The use of HBC for human use is problematic. HBC produced using whole oocysts is batch dependent and this may lead to the development of passive immune preparations which are not uniform in immunogenicity and potency. This generates a problem when these immune preparations are to be administered to human patients as such non-uniformity may result in failure of protection. In addition, it would be desirable to allow preparation of large amounts of antigen expressed in heterologous systems rather than to purify the oocyst.

Thus, there is a continuous need for immunogenic agents which are reasonably reproducible and have uniform and controllable immunogenicity and potency, which agents

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would be useful for the active and passive immunotherapy of cryptosporidiosis in both uncompromised and immunocompromised subjects, such as AIDS patients.

Additionally, there is a need to have available methods for reproducible expression of specific targets for Cryptosporidium antigens in large amounts, which antigens would provide a better immunogen. This approach requires that specific Cryptosporidium antigen genes are cloned and identified as potential candidates through their ability to elicit an antibody response that is immunoprotective. Before antibodies produced in this manner are tested in or administered to humans or animals, testing in in vitro assay of its inhibitory effect on invasion or intracellular development of the Cryptosporidium organism in cultured cells, and in vivo studies, would be desirable.

It is, therefore, a primary objective of this invention to provide polyclonal or monoclonal antibodies to be used for prophylaxis and treatment of cryptosporidiosis and to express a portion of the GP900 sequence/locus to provide target protein antigens allowing production of recombinant anti-Cryptosporidium vaccines and passive immune products.

All patents, patent applications and publications cited herein are hereby incorporated by reference.

SUMMARY OF THE INVENTION

One aspect of this invention concerns vaccines, antigens, antibodies, proteins, glycoproteins, DNAs and RNAs for prophylaxis and treatment of Cryptosporidium or Cryptosporidium infections.

Another aspect of this invention concerns a Cryptosporidium antigen comprised of an immunogenic protein without attached carbohydrates.

Another aspect of this invention concerns a Cryptosporidium antigen comprised of an immunogenic protein with attached carbohydrates.

Another aspect of this invention concerns polyclonal or monoclonal antibodies directed against the Cryptosporidium antigen.

Another aspect of this invention concerns DNA and RNA

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encoding or representing the Cryptosporidium antigen and fragments thereof.

Another aspect of this invention concerns polyclonal or monoclonal antibodies directed against invasive stages of cryptosporidial species capable of preventing and ameliorating the invasion of *Cryptosporidium* infection.

Another aspect of this invention concerns a synthetic or recombinant vaccine, useful for active immunization of animals and humans against Cryptosporidium infection.

Another aspect of this invention concerns a synthetic or recombinant protein useful for preparation of passive immune products for treatment of established infections.

Another aspect of this invention concerns a synthetic or recombinant DNA vaccine, capable of endogenous development of an inhibitory amount of anti-Cryptosporidium parvum antibodies.

Another aspect of this invention concerns a synthetic or recombinant RNA vaccine, capable of endogenous development of an inhibitory amount of anti-Cryptosporidium parvum antibodies.

Another aspect of this invention concerns an amino acid sequence (1721 aa) of a protein portion of GP900, a >900 kDa glycoprotein of sporozoites and merozoites, and its amino acid and size variants.

Another aspect of this invention concerns the DNA sequence of 5319 nucleotides encoding a protein portion of GP900, its nucleotide and size variants and its upstream (5') protein coding and regulatory elements.

Another aspect of this invention concerns the RNA sequence determined by the DNA sequence of GP900 and its nucleotide and size variants including the polyadenylation sequence.

Another aspect of this invention concerns an amino acid sequence (503 aa) of a protein portion of P68, a 50-100 kDa glycoprotein of sporozoites and merozoites, and its amino acid and size variants.

Another aspect of this invention concerns the DNA sequence of 2380 nucleotides encoding a protein portion of P68, its nucleotide and size variants and its upstream (5')

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protein coding and regulatory elements.

Another aspect of this invention concerns the RNA sequence determined by the DNA sequence of P68 and its nucleotide and size variants including the polyadenylation sequence.

Another aspect of this invention concerns the group of GP900 or P68 recombinant or expressed protein, or glycoprotein targets of antibodies which inhibit infection, invasion, or adhesion.

Another aspect of this invention concerns a method for prophylaxis and treatment of Cryptosporidium or Cryptosporidium infections using vaccines, antibodies, proteins, glycoproteins, DNAs and RNAs of the invention.

Another aspect of this invention concerns a method of prophylaxis, treatment, inhibition or retardation of a Cryptosporidium infection, comprising administering to a subject in need of such treatment an amount of anti-Cryptosporidium polyclonal or monoclonal antibodies, prophylactically or therapeutically effective, to provide immunity against infection or treatment for the disease.

Another aspect of this invention concerns a method of prophylaxis, treatment, retardation, or inhibition of Cryptosporidium infection, comprising administering to a subject in need of such treatment, a vaccine containing the polypeptide or glycoprotein of this invention or its DNA or RNA, capable of endogenous stimulation of the production of an inhibitory amount of anti-Cryptosporidium antibodies.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is an immunoblot of Cryptosporidium parvum oocyst/sporozoite proteins showing detection of the >900 sporozoite protein with monoclonal and polyclonal antibodies to GP900.

Figure 2 is the immunoprecipitation of ¹²⁵I surface label *Cryptosporidium parvum* sporozoite proteins using monoclonal and polyclonal antibodies to GP900.

Figure 3 is the MAb 7B3 indirect immunofluorescence detection of GP900 present on the surface and shed from the surface of a motile sporozoite.

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Figure 4 is the immunoblot of Cryptosporidium parvum N-deglycosylated sporozoite/oocyst proteins using monoclonal antibodies to GP900.

Figure 5 depicts the electronmicrographic localization of GP900.

Figure 6 is the genomic Southern analysis of the GP900 gene fragment.

Figure 7 shows an immunoblot using antibodies to recombinant GP900 proteins.

Figure 8 is a graphical illustration of the inhibition of invasion and intracellular development of MDCK cells in vitro by antibodies to GP900 recombinant proteins.

Figure 9 is a graphical illustration of the dose-dependent inhibition of sporozoites invasion/intracellular development in MDCK cells in vitro by affinity purified anti-S34 antibody.

Figure 10 is a graphical illustration of the inhibition of parasite burden in vivo in neonatal mice challenged with Cryptosporidium and treated with oral anti-recombinant GP900 antibodies.

Figure 11 is an immunoblot of Cryptosporidium parvum occyst/sporozoite proteins using polyclonal antibodies to P68.

Figure 12 is a graphical illustration of the inhibition of invasion and intracellular development of MDCK cells *in vitro* by antibodies to P68 recombinant protein.

Figure 13 is a graphical illustration of the inhibition of parasite burden <u>in vivo</u> in neonatal mice challenged with *Cryptosporidium* and treated with oral antirecombinant P68 antibodies.

DEFINITIONS

As used herein:

"GP900" means a high molecular weight protein represented by 1721 amino acids and identified as SEQ ID NO: 5 of Mr greater than 900 kilodaltons (kDa) which may have an attached glycoprotein, said GP900 detected at the surface of sporozoites or merozoites. GP900 is the target of antibodies which inhibit infection, invasion or adhesion

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of Cryptosporidium.

"P68" means an apical protein of sporozoites or merozoites represented by 503 amino acids and identified as SEQ ID NO: 6 of Mr between approximately 50 and 100 kilodaltons which is a target of antibodies which inhibit infection, invasion or adhesion of Cryptosporidium.

The "structure" or "structural characteristics" of GP900 defines a protein, glycoprotein, DNA and RNA encoding the GP900 protein and includes all structural variations, mutations and fragments exhibiting the same function.

The "functionality" or "functional characteristics" of GP900 is defined by the interaction of antibodies to GP900 and structural variants described, such that the antibody inhibits infection, invasion or adhesion of Cryptosporidium.

"T-cell epitope" means targets which stimulate or elicit T cells or T cell mediated immune responses.

"Cell mediated immune responses" means responses stimulated or elicited by interaction of T cell epitope or T cell.

"The gene" or "genes encoding GP900" means DNA encoding a portion or all of the GP900 protein. One or more of these portions with or without carbohydrates attached include the targets of GP900 antibodies known as T-cell epitopes.

The "structure" or "structural characteristics" of P68 defines a protein, DNA and RNA encoding the P68 protein and includes all structural variations, mutations and fragments exhibiting the same function.

The "functionality" or "functional characteristics" of P68 is defined as the interaction of antibodies to P68 and structural variants described, such that the antibody inhibits infection, invasion or adhesion of Cryptosporidium.

35 "The gene" or "genes encoding P68" means DNA encoding a portion or all of the P68 protein. One or more of these portions include the targets of P68 antibodies known as T-cell epitopes.

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"Sporozoites or merozoites" means any life stage which may invade host cells and any variant or mutant of said sporozoites or merozoites.

"Antibodies" means proteins which structurally interact with the target antigen and are produced when the antigen is introduced into an animal, such that they stimulate the immune system. The term also includes antibodies produced in vitro, such as antibodies produced by hybridoma cell cultures and chimeric proteins, as well as hybridoma cells and chimeric constructs introduced into the host to provide an in vivo antibody.

"Antibodies to GP900" means proteins which structurally interact with the target antigen GP900 and inhibit infection, invasion or adhesion of the sporozoites or merozoites to the host cell.

"Antibodies to P68" means proteins which structurally interact with the target antigen P68 and inhibit infection, invasion or adhesion of the sporozoites or merozoites to the host cell.

"Monoclonal antibodies" means the monovalent antibodies produced by B cells fused to immortalized cells producing specific antibody to GP900 or P68.

"Polyclonal antibodies" means antibodies directed at GP900 or P68 which are not monovalent and are the products of multiple B cells in character.

"Carbohydrate" or "carbohydrate moiety" means any Nor O-linked carbohydrate or portion thereof, which is covalently linked to the protein of GP900 or P68.

"Target antigen" means protein or carbohydrate moiety attached to protein including variants defined by differential glycosylation and conformational change.

"Differential glycosylation" means glycoproteins which vary in the carbohydrate moieties attached to the protein backbone as a function of factors other than the sequence of the protein backbone.

"Conformational change" means change in the shape of the protein or the glycoprotein as a result of changes in the carbohydrate moieties bound to it and/or changes in the protein sequence. . WO 98/06430

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"GP900 antigen" means a protein with or without a carbohydrate attached thereto which defines the capacity of Cryptosporidium sporozoites and merozoites to infect host cells.

"P68 antigen" means a protein with or without a carbohydrate attached thereto which defines the capacity of Cryptosporidium sporozoites and merozoites to infect host cells.

"GP900 DNA" means the sequence of 5319 nucleotides identified as SEQ ID NO.: 2 which encodes an amino acid portion of the protein sequence of GP900 protein (SEQ ID NO: 5) and any variant, 5' extension, mutation and fragment thereof, which corresponds to genes encoding the antigen.

"GP900 RNA" means the RNA sequence corresponding to 5164 bp of DNA sequence (SEQ ID NO. 1) which encodes the protein sequence of GP900 protein (SEQ ID NO: 5) and any 5' extension, variant, mutation and fragment thereof.

"P68 DNA" means the sequence of 1509 nucleotides identified as SEQ ID NO: 3 which encodes the protein sequence (SEQ ID NO: 6) of P68 protein and any 5' extension, variant, mutation and fragment thereof which corresponds to genes encoding the antigen.

"P68 RNA" means the RNA sequence corresponding to the 1509 nucleotides identified as SEQ ID NO: 2 which encodes the protein sequence of P68 protein (SEQ ID NO: 6) and any 5' extension, variant, mutation and fragment thereof.

"Vaccine" means a protein, recombinant protein, DNA or RNA from GP900 or P68 which upon introduction into a host, is able to provoke an immune response including but not limited to the production of antibodies, cytokines and other cellular responses.

"Prevention or prophylaxis" means the passive or active immunization with antibodies or vaccines of the invention such that disease or infection does not occur.

"Treatment" means therapeutic use of any protein, glycoprotein or antibody to inhibit existing infection in a host.

"Host" means a human or animal, including birds and cattle.

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"Regulatory elements" means nucleotide sequences which control the expression of genes they regulate, typically by interaction with other macromolecular species such as proteins.

"Active immunity to infection" means the ability of an organism to produce specific responses such as production of cytokines, lymphokines, antibodies or other substances, or cellular capacity to inhibit or retard infection in response to a contact with an antigen.

"Passive immunity to infection" means the transfer to a host of the specific antibodies or other substances or cells capable of inhibiting or retarding infection.

"Cryptosporidium species" means any organism belonging to the genus Cryptosporidium, such as, for example, Cryptosporidium parvum or Cryptosporidium muris, but also includes currently less well characterized other organisms such as, for example, Cyclospora and it is also meant to include apicomplexan parasites which invade the gastrointestinal tract, such as Eimeria. Cryptosporidium species comprise Apicomlexan parasites which primarily invade cells of the gastrointestinal tract and cause disease in a susceptible host.

"Recombinant vaccines" means 1) protein segments produced from recombinant DNA or RNA in vitro and then introduced in vivo; and 2) RNA or DNA introduced in vivo and capable of producing recombinant protein in situ. This term includes all vaccines other than those biologically derived. "Biologically derived vaccines" means a protein or glycoprotein generated in the organism of origin.

"REA" means recombinant eluted antibodies.

DETAILED DESCRIPTION OF THE INVENTION

This invention relates to vaccines, antibodies, proteins, glycoproteins, DNAs and RNAs useful for prophylaxis and treatment of infections caused by any Cryptosporidium organism or any organism belonging to the Cryptosporidium species.

More specifically, the invention concerns: identification and isolation of *Cryptosporidium* antigens comprised of a protein or polypeptide with or without a

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carbohydrate attached thereto; identification of the DNA of Cryptosporidium antigen gene within the sequencing DNA encoding Cryptosporidium antigens; of the locus encoding the expressing portions Cryptosporidium antigens; and using the expressed antigens to prepare vaccines or polyclonal or monoclonal antibodies.

I. Cryptosporidium Antigens

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Cryptosporidium organisms and particularly Cryptosporidium parvum are coccidian parasites of the gastrointestinal tract that cause a clinical syndrome of diarrhea for which there is currently no effective treatment. Infectivity of Cryptosporidium is mediated by a specific protein or polypeptide antigens of sporozoites or merozoites, the infective forms of Cryptosporidium.

Two antigen proteins, designated GP900 and P68 were They were partially sequenced at the DNA identified. The 3' sequence and 3' flanking regions for GP900 level. Two GP900 DNA sequences and two and P68 are documented. SEO ID NO: 1 DNA sequences were established. comprising 5164 bp encodes a portion of GP900. SEO ID NO: 2 comprising 5319 bp encodes a portion of GP900 and SEQ ID NO: 3 comprising includes the 3' flanking region. 1509 b; encodes a portion of P68. SEQ ID NO: 4 comprising 2380 bp encodes a portion of P68 and includes 3' flanking The deduced partial sequences of encoded proteins GP900 (SEQ ID NO: 5) and P68 (SEQ ID NO: 6) were established.

The DNA encoding Cryptosporidium antigen can be coupled to Cryptosporidium DNA encoding regulatory elements located downstream or upstream or on another chromosome in the Cryptosporidium genome. These operably coupled DNA segments are able to bind selectively and specifically to Cryptosporidium molecules, such as proteins.

Expressed portions of the loci encoding GP900 and P68, are targets of polyclonal and monoclonal antibodies able to inhibit invasion/intracellular development in vitro and in vivo. The expression, identification and isolation of these recombinant proteins allows production of recombinant vaccines for active immunization of animals and humans

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against cryptosporidiosis as well as passive immune products for prevention and treatment of an established infection.

During the development of this invention, it has been shown and described in Infect. Immun., 60:2343 (1992), 60:5132 (1992), and 61:4079 (1993), that a Cryptosporidium parvum expression library clone S34 encoded a portion of a protein larger than 900 kDa, recognized by hyperimmune bovine colostrum (HBC), which has been designated GP900.

The GP900 protein is highly abundant and is easily visualized by Coomassie blue staining of proteins on SDSpolyacrylamide gels (SDS-PAGE). Furthermore, it is known to be Triton X-100 soluble and N-glycosylated.

This protein has been localized at the anterior portion of the sporozoite and merozoite immunofluorescence microscopy. Specifically, the protein has been detected in micronemes of these invasive stages by immunoelectronmicroscopy and has been shown to be accessible to surface radioiodination with 125I.

Monoclonal antibodies, which are specific for GP900, have been made according to Example 2. Three of six antibodies, namely 10C6, 7B3, and E6, made from a single fusion event in which the immunogen was an containing sporozoites, were specific to GP900, suggesting that GP900 is a highly immunogenic molecule of sporozoites. 25 Three of eight antibodies, namely M2, M15 and M24 made from a second fusion event, in which the immunogen consisted of meronts, were also specific to GP900, suggesting that GP900 is a highly immunogenic molecule of merozoites.

30 The second Cryptosporidium antigen is a smaller protein identified as P68.

GP900 Protein, Glycoprotein, Recombinant Protein 1. and DNA/RNA

Cryptosporidium Antigen Protein GP900 A.

Cryptosporidium antigen GP900 is a high molecular 35 weight glycoprotein of a Mr greater than 900 kilodaltons The GP900 protein was detected in micronemes of developing merozoites and sporozoites. It is present on the surface of the sporozoites and is shed from the

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sporozoite surface in vivo in host cells. When deglycosylated, the GP900 core protein has a variable molecular weight of approximately 150-250 kDa. The GP900 protein has been identified as a target of anti-GP900 antibodies which inhibit Cryptosporidium infection, invasion or adhesion.

The GP900 protein sequence containing 1721 amino acids is seen in SEQ ID NO: 5. DNA encoding the GP900 protein contains 5164 base pairs and its sequence is depicted in SEQ ID NO: 1.

Cryptosporidium parvum was identified and isolated oocysts the Iowa and AUCP-1 of isolates Cryptosporidium parvum passaged through neonatal calves. as described in Example 1. Occysts containing sporozoites were solubilized, resolved by SDS-PAGE and subjected to immunoblotting, according to Infect. Immun., 60:5132 Proteins which are targets of oocyst/sporozoite antibody were visualized after incubation with the primary antibody by enzyme-linked immunosorbent assay (ELISA) or with 125I labeled Protein A followed by autoradiography.

Protective HBC Ig antibodies were, during the development of this invention, found to react predominantly with two proteins above 200 kDa in a Western blot of solubilized occyst. The first protein had a size of 250 kDa. The second protein was >900 kDa and comigrated with GP900. In an attempt to determine whether the 250 kDa protein is a component of GP900, polyclonal antibodies against SDS solubilized GP900 were prepared.

Identification of the GP900 protein from the cocyst of Iowa and AUCP-1 isolates is illustrated in Figures 1 and 2.

Visualized Cryptosporidium antigen proteins were surface radioiodinated and immunoprecipitated using the method described in Infect. Immun., 61:4079 (1993).

Triton X100 soluble GP900 of sporozoites was used to prepare polyclonal anti-GP900 antibody to clone the gene for GP900. To this end, the soluble fraction of GP900 was immunoprecipitated with monoclonal antibody (MAb) 10C6, and a >900 kDa molecular weight species was identified and

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excised, and used for immunization of mice for production of the anti-GP900 antibody. Polyclonal antibodies prepared against SDS solubilized GP900 and MAb 10C6, which were previously shown to detect GP900, were used to probe a Western blot, as seen in Figure 1, and to immunoprecipitate sporozoite surface labeled proteins, as seen in Figure 2.

Figure 1 shows a immunoblot of Cryptosporidium parvum oocyst/sporozoite proteins of the AUCP-1 isolate separated by SDS-PAGE. Lane 1 shows the MAb 10C6 culture supernatant. Lane 2 shows the polyclonal anti-GP900 in 1:5000 dilution.

As seen in Figure 1, a single molecular species, GP900, was identified at ~900 kDa by both monoclonal and polyclonal antibodies. Cross-immunoprecipitation studies confirmed that the same, approximately 900 kDa size, protein was seen by both antibodies. At prolonged periods of detection, a less prominent ladder of bands between the 200 and 92 kDa markers was observed.

Figure 2 shows immunoprecipitation of ¹²⁵I radiolabeled Cryptosporidium parvum sporozoite surface proteins of the AUCP-1 isolate separated by 5-15% SDS-PAGE. Lane 1 shows radiolabeled Cryptosporidium parvum sporozoite surface protein control (10⁷ sporozoites/lane). Lane 2 shows radiolabelled Cryptosporidium parvum sporozoite surface proteins immunoprecipitated with monoclonal MAb 10C6. Lane 3 shows radiolabelled Cryptosporidium parvum sporozoite surface proteins immunoprecipitated with polyclonal anti-GP900.

Immunoprecipitation of ¹²⁵I labeled sporozoites with polyclonal anti-GP900 and monoclonal 10C6 antibodies revealed that polyclonal anti-GP900 only detects one protein, GP900, while monoclonal 10C6 additionally detects a protein of 250 kDa. These data suggest that MAb 10C6 detects a shared epitope on two surface accessible proteins, GP900 and a protein of Mr 250,000. However, the 250 kDa protein also could be a precursor or processed form of GP900 and as such is a part of the invention.

These results show that polyclonal anti-GP900 antibody is a more specific detection reagent for GP900 than monoclonal Ab10C6 by Western blot and by

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immunoprecipitation of surface proteins. This confirms that polyclonal anti-GP900 antibody is an appropriate antibody for GP900 localization experiments and for detection of clones in a *Cryptosporidium* expression library.

Prominent sporozoite surface proteins of other Apicomplexan parasites, for example, the circumsporozoite protein of the *Plasmodium* species, which contains the binding ligand for adhesion of the malaria sporozoite to its host cell, the hepatocyte, are known to be shed from the surface of sporozoites in vivo.

In order to determine whether GP900 similarly was shed from the surface of *Cryptosporidium* sporozoites, living sporozoites were allowed to glide on poly-L-lysine coated slides. Results are shown in Figure 3.

Figure 3 shows the indirect immunofluorescence detection of GP900 with MAb 7B3 after fixation of the sporozoites with formaldehyde. MAb 7B3 was used because it was previously shown to detect only GP900 on immunoblots of Cryptosporidium sporozoite proteins. Figure 3 shows that GP900 is present around the living sporozoite and is shed from the posterior aspect of living sporozoites as the sporozoites glide.

In order to show that GP900 is a glycoprotein, N-linked carbohydrate was enzymatically removed from Cryptosporidium parvum oocyst/sporozoite proteins and the remaining protein was as separated by SDS-PAGE and detected with MAb 10C6 by an immunoblot. Results are seen in Figure 4.

Figure 4 indicates that deglycosylation removes reactivity with MAb 10C6 (1), a GP900 reactive MAb, and generates new reactivities (2) which exhibit a ladder-like pattern between markers having apparent molecular weight 97 and 200. The estimated Mr of those molecular species in this figure are 150,000-180,000. This data is consistent with the removal of N-linked carbohydrate moieties from the GP900 protein backbone and with the appearance of the protein backbone alone or with incomplete removal of all N-linked carbohydrates or O-linked sugars. It is also

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consistent with potential size polymorphism in the GP900 core protein or combination of these events. From this data the protein core of GP900 would be predicted to have a Mr of 150,000-180,000.

An anti-GP900 polyclonal antibody, affinity purified from occyst/sporozoite antibodies on the protein expressed by a \(\lambda\)gtll clone S34 using S34 recombinant eluted antibody (S34 REA), also detected a Mr greater than 900,000 protein in intact sporozoites and a ladder of proteins with the smallest at about Mr 150,000 after N-deglycosylation. Cross immunoprecipitation experiments showed that these three reagents, S34 REA, and anti-GP900 polyclonal and monoclonal MAb 10C6 antibodies, detected the Mr >900,000 protein indicating that the S34 clone encodes a portion of the core protein which when glycosylated corresponds to the GP900 protein.

In order to determine the subcellular localization of GP900, anti-GP900 mouse ascites were assayed on LR White Electronmicrograph sections of ileum from experimentally infected rats that contained all developmental stages of the parasite. Results are seen in Figure 5.

In Figure 5, the micronemes of merozoites (within a developing schizont) appear as stacked plate-like radiolucent structures in which the gold particles of the antibody, used to localize GP900 polyclonal antibodies, are concentrated. GP 900 was also seen in sporozoites within oocysts (data not shown). The rhoptries and dense granules were not labeled. No surface labeling sporozoites and merozoites was observed. No gold particles were detected in the parasitophorous vacuole or over the vacuolar wall. No antigen could be detected in host-cell cytoplasm.

B. <u>GP900 Gene Cloning/Sequencing and Genomic Southern</u> Analysis

The GP900 gene of Cryptosporidium parvum was isolated from a naturally infected neonatal calf (NINC) isolate. DNA from calves was used to prepare a \(\lambda\gamma\text{t11}\) expression library containing clones with an open reading frame for GP900 which is 5164 bp. The sequence of this open reading

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frame was determined in the following way.

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Clone S34 was previously determined to encode a portion of a much larger protein, GP900, using S34 REA. A second GP900 clone, Ag4, reacted with polyclonal anti-GP900 and a MAb to GP900 but the Ag4 and S34 insert DNAs did not cross-hybridize with one another. Upon sequencing, the clones were found to contain distinctly different sequences with no overlap. The inserts were used to double screen the λ genomic library to determine if a clone encoding both could be identified indicating that Ag4 and S34 were Clone DB8, which collinear portions of the same gene. hybridized to both S34 and Ag4, contained a single open reading frame containing both the S34 and the PCR amplification products generated from the seguences. 5' and 3' terminal sequences of DB8 were used to identify clones, 95-18 and 93-14, respectively. When added to the DB8 sequence, the sequence generated from these clones comprised the open reading frame which is still open at the 5' end and 3' flanking noncoding region.

Genomic Southern analysis was undertaken to determine if GP900 was encoded by a single gene and if so whether this gene was polymorphic in the three isolates for which the locus was available, namely the NINC isolate and the Iowa and AUCP-1 isolates. DB8 DNA from NINC isolate was used as a probe. The sequence of DB8 contains no EcoRI, Bgl II or Hind III sites but contains many (10) Hind I sites. The larger restriction fragment has a size of 1146 bp and include parts of domains 1 and 2. The short restriction fragment is 741 bp and covers half of domain 3 and all of domain 4. Results are seen in Figure 6.

Figure 6 is a genomic Southern analysis of the GP900 gene locus. The Southern blot was hybridized with the pDB8 insert. Lanes 1-4 shows the Iowa isolate DNA. Lane 5 shows the AUCP-1 isolate DNA. Lane 1 shows EcoR I digestion. Lane 2 shows Bgl II digestion. Lane 3 shows Hind III digestion. Lanes 4 and 5 show Hind I digestion. No difference in restriction pattern was seen between the Iowa and AUCP-1 isolates in the Hind I digestion lanes.

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Figure 6 shows that the DB8 probe of the NINC Cryptosporidium isolate hybridizes to a single DNA fragment in EcoRI, Bgl II and Hind III digests of the Iowa strand, indicating that GP900 is encoded by a single prominent gene. In lanes 4 and 5 the probe hybridizes to two 5 fragments in the Iowa and AUCP-1 isolate DNA, which are of approximately the same size as the two largest fragments encompassing the two polythreonine regions of the DB8 These data indicate that gross GP900 gene probe. rearrangements have not occurred in the three different isolates studied. This observation is further confirmed by the fact that both the Iowa and AUCP isolates produce a large 900 kDa protein which reacts with the polyclonal antibodies to GP900 initially prepared against the AUCP-1 15 isolate.

Structure of the GP900 Gene and its Encoded Protein C. Sequences identified as SEQ ID NO: 1 and 2 nucleotide sequences of the GP900 gene fragment. The sequence identified as SEQ ID NO: 5 is the corresponding protein.

The GP900 open reading frame encodes two mucin-like polythreonine domains.

Domains 1 and 3 of the protein are cysteine rich domains whereas domains 2 and 4 are mucin-like domains containing large numbers of threonines.

Domain 1 contains 5 cysteine residues. Domain 3 has 6 cysteines. Neither domain is highly homologous to any known sequence in GenBank or Swiss Protein Bank.

Domain 2 is composed of 94% threonine residues including an unusual stretch of 112 uninterrupted threonines. Domain 4 is composed of 56% threonine residues. Both domains also contain repeats of the sequence lysine-lysine-proline or lysine-proline. When the deduced protein sequence was analyzed by searches of the GenBank and Swiss Protein Bank, the greatest similarities were found between the threonine-rich regions of GP900 and other glycoproteins with either proven or putative 0-linked glycosylation.

GP900 is both N- and O-glycosylated. GP900 has been shown to be susceptible to treatment with N-glycosidase F (N-glycanase) which cleaves high mannose and complex structures (Figure 4).

The presence of abundant cysteines on a surface protein of Cryptosporidium which is functionally homologous to the circumsporozoite protein of malaria strongly suggests that these cysteines participate in binding phenomena and may comprise new binding motifs. Numerous apicomplexan parasite proteins, such as plasmodium, CSP, Duffy binding protein, EBA and PFEMPI have binding domains which contain cysteine rich regions. N or O linked carbohydrate moieties may also participate in binding to adjacent cells.

15 D. Production of GP900 Recombinant Proteins

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In order to prepare reagents for specific portions of GP900 to assay their effects on sporozoite adhesion, invasion and intracellular development in vitro and infection in vivo, polyclonal antibodies were made to purified wild type β -galactosidase, Ag4- β -galactosidase and S34- β -galactosidase fusion proteins according to Example 6.

In order to further define these antibodies by removing the reactivity to β -galactosidase and to concentrate them, affinity purified antibodies to the Ag4 and S34 portions of their fusion proteins were prepared according to Example 6. These various antibody preparations were used to probe an immunoblot of proteins from Cryptosporidium parvum occysts/sporozoites. Results are shown in Figure 7.

Figure 7 is an immunoblot of proteins obtained from Cryptosporidium parvum oocysts or sporozoites. Marker size in kDa is indicated. Lane 1 is the S34 antigen probed with pre-immune rabbit serum. Lanes 2-4 are the serum of rabbit immunized with S34 antigen. Lane 2 is probed with anti-S34 antibody. Lane 3 is probed with the polyclonal anti-Ag4 antibody. Lane 4 is probed with the anti-Ag4 affinity purified (A.P.) polyclonal antibody.

Figure 7, lane 1, shows that the pre-immune serum from the rabbit which received the S34 antigen is mildly

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reactive to two proteins of Cryptosporidium parvum. After immunization with the S34 antigen (lanes 2-4), the antisera react with a whole variety of proteins including GP900, a ladder of proteins ranging in size from 150 to 250 kDa, and several different proteins of lower molecular weight. Since the S34 sequence carries the poly-threonine repeats, it would seem that the antibody which recognizes these repeats will also recognize other proteins with this repeated motif and that the multiple bands represent such cross reactions. However, the results point toward another interpretation. The polyclonal antibody directed against Ag4, which does not carry poly-threonine repeats, and the affinity purified Ag4 antibody, recognize GP900 as well as the ladder of proteins between 150-250 kDa, suggesting that the ladder protein represents the core protein of GP900, not cross-reacting proteins.

E. In Vitro and In Vivo Assessment of Activity of Anti-GP900 and Anti-Recombinant GP900 Antibodies

In order to determine whether native or recombinantly produced antibodies in fact inhibit Cryptosporidium infection and would, therefore, be viable reagents for provoking active or providing passive immunity, or be useful for therapeutic purposes, in vitro and in vivo assessments of the effect of native or recombinant antibodies raised against GP900 antigen were investigated.

Antibody mediated inhibition of invasion and intracellular development was studied in Madin Darby Canine Kidney (MDCK) cells.

For these studies, MDCK cell monolayers were infected with Cryptosporidium parvum oocysts of the Iowa isolate in the presence of control reagents or immune sera, and colostrum was directed against a series of Cryptosporidium parvum oocyst antigens. Antisera and HBC Ig/sham HBC Ig were diluted 1:40 in cell culture media, such as RPMI, except for affinity purified anti-Ag4 which was diluted 1:16. Affinity-purified anti-Ag4 was assayed at a final protein concentration of 75 μ g/ml by the Bradford technique. The protein concentration of HBC Ig 40529 at a

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1:40 dilution was 800 μ g/ml. Results are shown on Figure 8.

Figure 8 is a graph showing Cryptosporidium parvum invasion into MDCK cells which were not treated (RPMI control = Bar 1) or were treated with fetal calf serum (bar 2), with sham HBC serum (bar 3), with HBC Ig 40529 serum (bar 4), with anti- β -galactosidase serum (bar 5), with anti-GP900 serum (bar 6), with anti-S34 serum (bar 7), with anti-Ag4 serum (bar 8) or with anti-Ag4 affinity purified serum (bar 9). Responses are expressed in percent of invasion.

in Figure As seen 8, inhibition of invasion/intracellular development was observed with antisera raised against protein epitopes of Aq4 (8) and S34 (7) expressed as β -galactosidase fusion proteins. antibody raised against wild type β -galactosidase (4) did not confer protection. Negative experimental controls for the inhibition assay included RPMI containing no additives (1), RPMI containing 5% fetal calf serum (2), or sham hyperimmune bovine colostrum (SHAM-HBC) (3), collected from COWS immunized with herd vaccines but not with Cryptosporidium. Recombinant S-34-glutathione-stransferase fusion protein (S34-GST) at 100 nM and 1 μ M preincubated with 1:40 anti-S34-galactosidase (bar 4 shows 1 μ M) abolished the inhibitory activity of the antibody verifying that the specificity of the inhibition is conferred by the S34 protein sequence and the antibody to it.

Figure 8 clearly shows that the absence of antibodies in controls or the presence of sham HBC anti- β -galactosidases and anti-GP900 antibodies did not provide protection against Cryptosporidium infectivity, invasion or adhesion. On the other hand, antibodies raised against S34 (bar 7) and against Ag4 (bar 8 and bar 9), whether affinity purified (bar 9) or not (bar 8), provided good protection against Cryptosporidium infection. Affinity purified polyclonal antibody (bar 9) was the most active in this system and almost equal in activity to HBC immunoglobulin.

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In order to determine whether the inhibitory response depends on the dose of the antibody, dose response curve of the affinity purified S34 antibodies at concentrations of 10 (bar 5), 50 (bar 6), 100 (bar 7) and 500 (bar 8) μ g/ml, was determined and compared to controls represented by untreated oocysts (bar 1), S34- β -galactosidase antibodies (bar 2), HBC Ig (bar 3) and anti-oocyst antibodies (bar 4). Response is expressed in parasite-to-host nuclei ratio. Results are seen in Figure 9.

As seen in Figure 9, 50 μg/ml of anti-S34 antibody provided more than 50% protection while 100 μg/ml and 500 μg/ml provided excellent to almost complete protection against invasion and intracellular development of Cryptosporidium sporozoites. These results further demonstrate the specific nature of the antibody/antigen reaction.

In order determine to whether the native recombinant antibodies raised against Cryptosporidium antigen GP900 or a fraction thereof are able to inhibit Cryptosporidium infection in vivo, the anti-S34-βgalactosidase and anti-S19- β -galactosidase antibodies were tested in a neonatal mice model. Results are seen in Figure 10.

Figure 10 shows the *in vivo* effect of antibodies to GP900 recombinants on shedding of oocysts by neonatal mice infected with *Cryptosporidium*.

Figure 10 is a graph representing the amount of excretion of Cryptosporidium oocysts per day in mice treated with phosphate buffered saline (bar 1); anti- β -galactosidase (bar 2); anti-Ag4- β -galactosidase (bar 3); anti-S34- β -galactosidase (bar 4); 1:5 HBC Ig 40529 (bar 5); and paromomycin (bar 6). As seen in Figure 10, anti-S34 (bar 4) reduced the oocysts shed by about 50% relative to control PBS (bar 1) and anti- β -galactosidase antibody (bar 2). Although crude antisera was used, antibody to S34 β -galactosidase inhibited shedding by about 50% relative to control treated with PBS and anti- β -galactosidase antibody. The inhibition was superior to the inhibition conferred by a 1:5 dilution of HBC Ig 40529 (bar 5), the positive

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control antibody which had previously been shown to prevent cryptosporidial disease in calves challenged with Cryptosporidium (Infect. Immun., 61:4079-4084 (1993)).

From the results obtained in these experiments, it is clear that clone S34 encodes a Cryptosporidium antigen and that the antibodies specifically raised against this antigen are able to inhibit the Cryptosporidium infection in vivo.

In order to determine whether GP 900, like the is circumsporozoite protein of malaria, an glycoprotein mediating the attachment of the sporozoite to a cell of GI origin, a paraformaldehyde fixed CaCO-2 cell adhesion assay was used to assess antibodies to β galactosidase, Ag4- β -galactosidase and S34- β -galactosidase, In this assay, the same as described in Example 13. magnitude of inhibition of adhesion of Cryptosporidium sporozoites to CaCO-2 cells (mean O.D. 50% of control in ELISA) with a 1:50 dilution of anti S34- β -galactosidase was conferred as was observed in the in vitro invasion and intracellular development assay in living MDCK cells by a 1:40 dilution of the same antibody as seen in Figure 8.

In addition, these results were comparable to those seen when a 1:100 dilution of anti-Cryptosporidium murine (48% inhibition) a polyclonal rabbit Cryptosporidium antiserum (inhibition 51%) were previously assayed in this system (data not shown). Similarly to the in vivo model, in this in vitro model, the anti-Ag4- β galactosidase also did not inhibit invasion and infection anti-534 inhibited development. However, invasion/intracellular development in living MDCK cells in vitro, adhesion in killed CaCO-2 cells in vitro and These results support the infection in vivo in mice. premise that a biological function is inhibited by anti-S34 antibodies in the in vitro and in vivo systems and that that function is adhesion. Additionally, these results show that antibodies to recombinant GP900 significantly with the inhibitory activity of HBC Ig 40529 and anti-Cryptosporidium antibodies from mouse and rabbit sources.

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The antigen has been deduced to have an amino acid sequence depicted by SEQ ID NO: 5 and was encoded by genomic DNA sequences depicted by SEQ ID NO: 1. Antibodies against the recombinant S34 protein are able to significantly inhibit Cryptosporidium infection in vitro and in vivo.

Consequently, the results described above indicate the usefullness of the anti-S34 antibody for both anti-Cryptosporidium prophylaxis and therapy of a human or animal host.

2. P68 Protein, Recombinant Protein and DNA/RNA

A. <u>Identification of the Cryptosporidium Antigen</u> designated P68

A Cryptosporidium antigen designated P68 is an apical protein of sporozoites and merozoites. The protein has a size of between about 50-100 kDa. The P68 protein consists of 503 amino acids and its amino acid sequence is depicted as SEQ ID NO: 6. The P68 protein is derived from the gene S19. The DNA sequences encoding the P68 protein are depicted as SEQ ID NO: 3 and 4.

B. <u>Cloning/Sequencing and Genomic Southern Analysis of</u> the <u>Gene for P68</u>

The purification and initial characterization of the S19 clone and the description of the restriction fragment genomic expression library from which it was isolated have been described in (Infect. Immun., 60:5132-5138 (1992)). A recombinant eluted antibody from the clone identified a dominant 68 kDa protein on Western blot (Figure 11) of occyst sporozoite proteins and was localized to the anterior end of the sporozoite by indirect fluorescent antibody analysis.

Figure 11 is an immunoblot of AUCP isolate oocyst/sporozoite proteins. Lane 1 was detected with polyclonal anti-sporozoite/oocyst antibodies which had been affinity purified on the S19 fusion protein (S19-REA). seen in Figure 11, an immunoblot with the antibody identified the protein of Mr less than 69 kDa marker, known as 68 kDa protein. Lane 2 is REA prepared on wild type β galactosidase as a negative control.

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The S19 insert was subcloned into Bluescript and sequenced as described in Example 17. The insert was used as a molecular probe to identify λ gtl1 expression library clones which extended 5' and 3' from S19. A 2380 bp locus was defined. The defined portion of the locus has 1509 bp of open reading frame which remains open at the 5' end.

C. Structure of the Gene and its Encoded Protein

The sequences of the P68 gene fragment are shown in SEQ ID NOs: 2 and 3. The sequence of the corresponding protein is given in SEQ ID NO: 6.

D. Production of P68 Recombinant Proteins

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Using essentially the same methods as described for GP900 and for clones S34 and Ag4, S19 was subcloned into the pGEX expression vector to yield the expression clone GST-S19, a recombinant protein fused to glutathione S transferase. Antibodies were raised to GST-S19 in two rabbits and to the native GST, according to methods described in Examples 4 and 5.

E. <u>In Vitro</u> and <u>In Vivo</u> Assessment of Activity of Anti20 <u>P68</u> Antibody

Antibodies were assayed in vitro and in vivo for inhibition of invasion and inhibition of infection, respectively, as described in Examples 11 and 14. Results are seen in Figures 12 and 13.

The graph seen in Figure 12 shows inhibition of invasion and intracellular development by antibodies to P68 expressed as parasites/MDCK cell nuclear ratio. All antibodies were at a dilution of 1:40 except FCS.

As seen in Figure 12, controls such as RPMI medium (bar 1), preimmune serum (bar 2), 5% FCS (bar 3), and anti-GST fusion protein (bar 4) and (bar 5) antibodies raised against native GST were not effective in inhibiting Cryptosporidium infection. The two anti-GST/S19 (bar 6) and (bar 7) antibodies raised against a recombinant fusion protein of S19-glutathione-s-transferase clone inhibited the Cryptosporidium invasion by 46% and 33% relative to control. Both were more inhibitory than an anti-occyst/sporozoite (bar 8) antibody made in rabbits. The anti-GST antibodies, pre-immune antibodies and 5% fetal

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calf serum (FCS), did not inhibit invasion and intracellular development.

Figure 13 is a graph showing the effect of anti-S19 antibody on cocyst excretion in vivo in a CD neonatal mouse model as described in Example 14. Figure 13 shows that antibodies to S19-GST raised in two rabbits (bar 3) and (bar 4) significantly decreased cocyst excretion as compared to animals treated with anti-GST antibody (bar 2) or without treatment (bar 1).

II. <u>Mucosal and Systemic Lymphoproliferative</u> Responses

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In order to evaluate the immune response to Cryptosporidium infection in HIV infected or in healthy individuals, mucosal and systemic lymphoproliferative responses were studied using recombinant GP900 stimulation of mucosal lymphocytes in the gastrointestinal mucosa of a rhesus macaque infected with SIV and Cryptosporidium which did not have the clinical disease. This model was established to determine what type of mucosal responses are correlated with resistance to the clinical disease.

The exact experimental conditions are described in Example 15. Briefly, a rhesus macaque mucosal immunity model was developed to evaluate the immune response to Cryptosporidium in normal and in HIV infected individuals. A mucosal lymphocyte proliferation experiment, results of which are shown and described in Tables 1 and 2, monitored ³H uptake of cultured lymphocytes harvested at necropsy from the duodenum/jejunum, ileum, and colon of a clinically well, (no weight loss, no diarrhea) SIV-infected macaque, experimentally Cryptosporidium infected rhesus with a low CD4 count of 250/mm³. The animal intermittently excreted oocysts. The experiment was designed to look at T cell stimulation and used concanavalin A (con A) as a T cell mitogen control.

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TABLE 1

PROLIFERATION OF GASTROINTESTINAL MNCs
BY SPECIFIC ANTIGEN

	A. Cell Origin	<u>Unstimulated</u> (mean cpm)	US standard deviation	Con A (mean cpm)	Con A standard deviation	<u>\$.1</u> .
10	Duo/Jej IEL	743	293	2602	130	*3,52
	Duo/Jej LPL	640	30.7	151	31	0.23
	Ileum IEL	834	32.8	4021	544	*4.8
	lleum LPL	633	102	175	45	0.27
	Colon IEL	1170	457	2355	39.5	*2.0
15	Colon LPL	556	26	272	14.8	0.48
	Spleen	8726	204	143,588	1810	*16.4
	B. Cell Origin	<u>Unstimulated</u> (mean cpm)	US standard deviation	Spz Ag (mean cpm)	Spz Ag standard deviation	<u>S.I.</u>
20	Duo/Jej IEL	333	55	296	75	0.88
	Duo/Jej LPL	644	40	304	103	0.47
	Ileum IEL	500	144	388	96	0.77
	Ileum LPL	627	130	286	50	0.45
	Colon IEL	285	9.5	300	1.1	1.05
25	Colon LPL	891	62.5	521	62	0.58
	C. <u>Cell Origin</u>	<u>Unstimulated</u> (mean cpm)	US standard deviation	S34 (mean cpm)	\$34 standard deviation	<u>s.i.</u>
	Duo/Jej IEL	333	55	185	49	0.55
30	Duo/Jej LPL	644	40	1902	574	*2.95
	licum IEL	500	144	201	49	0.4
	lieum LPL	627	130	4357	403	*6.9
	Colon IEL	285	9.5	497	121	1.7
	Colon LPL	891	62.5	14720	3003	*16.5
35						10.5
	D. Cells Origin	Unstimulated (mean cpm)	US standard deviation	S19 (mean cpm)	S19 standard deviation	<u> S.I.S.</u>
	Duo/Jej IEL	333	55	500	22.8	1.5
	Duo/Jej LPL	644	40	537	79	0.83
40	Ileum IEL	500	144	1016	134	*2.03
	lleum LPL	627	130	478	144	0.76
	Colon IEL	285	9.5	1347	394	*4.72
	Colon LPL	891	62.5	720	126	0.8

Table 1 shows the incorporation of 3H by unstimulated mononuclear cells (MNCs) as mean counts per well and the incorporation in the presence of the T cell mitogen concanavalin A, sporozoite antigen at 10 μ g/well, S34-GST antigen at 10 μ g/well or S19-GST antigen at 10 μ g/well. MNCs were evaluated as a function of site in GI tract that is duodenum/jejunum, ileum or colon, and compartment of mucosa (LPL = lamina propria lymphocytes; IEL = intraepithelial lymphocytes).

55 S.I. is the stimulation index, expressed as the ratio of the stimulated to unstimulated wells. Values of S.I.

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greater than 2-3 are considered evidence of significant stimulation.

Results were obtained in quadruplicate with single outlying values discarded.

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SUITABLE PROLIFERATION OF SPLEEN CELLS BY SPECIFIC ANTIGEN

10	Antigen	Unstimulated	US standard	Stimulated	S standard	<u>S.I.</u>
	Spz Ag	1619	428	2658	36	1.64
	S19	1619	428	3158	51	1.95
	S34	1619	428	1336	51	0.82
	GST	1619	428	1700	278	1.05
15	Con A	8726	204	143,588	1810	*16.4

Table 2 reports data for spleen MNCs, representative of the systemic immune system of the same animal. Data found during these experiments shows that although spleen cell (systemic lymphocytes) responded to the T cell mitogen with a significant proliferation, they did not respond to the sporozoite antigen (Spz Ag), S19-GST (S19), S34-GST (S34) or the control peptide GST, indicating that there was no systemic proliferative response in the face of known Cryptosporidium infection without disease.

Mucosal lymphocytes did respond to the recombinant antigens S34 and S19 with significant proliferative responses in a compartment specific manner, (S34 in LPL, S19 in IEL compartments) indicating active mucosal immune responses to these antigens in the face of SIV infection in a clinically healthy animal with intermittent shedding of Cryptosporidium.

Previous studies in rhesus macaques indicated that intraepithelial lymphocytes (IELS) are primarily T cells and lamina propria lymphocytes (LPLs) are predominantly B cells. Evaluation of the FAC cell marker profiles of stimulated cells in each compartment of animals like this is expected to shed light on the type of mucosal immune response which is protective in rhesus macaques. Germane to the current application, the observation that both S19 and S34 stimulated MNCs at concentrations of antigens which did not lead to proliferation in the presence of whole sporozoite antigen indicate that both antigens stimulated

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the mucosa of this immunocompromised animal and might be viable immunogens for active immunization of immunocompromised animals.

These results suggest that GP900 and P68 epitopes contained in S34 and S19, respectively, stimulate lymphoproliferation of mucosal PMNs.

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second rhesus macaque that was SIV and Cryptosporidium infected remained clinically well developed diarrhea and was excreting Cryptosporidium as her blood CD4 count fell below 100/mm3. Lymphoproliferative responses to sporozoite antigens, S34-GST, S19-GST, GST and concanavalin A, could not be detected in the blood, spleen, or either compartment of the gastrointestinal mucosa. However, ELISPOT analysis, in which B cells secreting antibodies to specific antigens are detected, showed that some LPLs cells still retained the ability to secrete antibodies to S34-GST and sporozoite antigens assayed at 5 x the S34-GST concentration, but not to GST.

This indicates that the proliferative response to S34
20 and sporozoite antigen was lost at a time when the animal
was becoming clinically ill and still had B cells which
recognized GP900. These findings suggest that the
cytokines necessary for a vigorous B cell response to
foreign antigen had been lost.

25 These results. when taken together with the observations from the first animal, suggest that Cryptosporidium infection may cause chronic diarrhea when the mucosal proliferative response to cryptosporidial antigens ceases, with a fall in production of the antibody in the lamina propria to specific Cryptosporidium antigens 30 to subprotective levels.

III. In vitro Inhibition of Cryptosporidium parvum Infection

in vitro inhibition of the invasion intracellular development of Cryptosporidium described in 35 Example 11 for GP900 and in Example 18 for P68 protein was shown to occur as a function of anti-Cryptosporidium titer. This was evidenced by its correlation with the corresponding immunoglobulin concentration in protective

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colostrum (HBC), and by the lack of biological activity of SHAM colostrum (SHAM-HBC). In a supportive experiment described in Example 13, HBC Ig was also shown to significantly inhibit Cryptosporidium parvum adhesion in the CaCO-2 cell line, thus providing a potential mechanism for inhibition of invasion and infection.

The in vitro ability of HBC to prevent Cryptosporidium infectivity was shown to be mediated only by specific anti-Cryptosporidium antibodies eluted from Cryptosporidium. Elution from fetal calf serum and SHAM-colostrum did not produce inhibition of the infection.

The studies conducted in support of this invention show that antibodies to specific Cryptosporidium antigens are also responsible for the in vitro effect of the HBC Ig fraction. Additionally, the inhibition in the in vitro assay correlates well with the effect of HBC and HBC Ig in vivo. These results validate the in vitro MDCK cell model as a model for detecting antibodies which were found to be protective in vivo.

Antibodies raised to the fusion proteins of several of the antigens of the invention were found significantly inhibitory in the *in vitro* MDCK cell model. This finding was then confirmed by similar results obtained with another epithelial cell line, Madin Darby Bovine Kidney (MDBK) cells. These *in vitro* results indicated that these antibodies by correlation of *in vitro* results to *in vivo* systems, would be protective *in vivo*. This was later confirmed in the *in vivo* studies.

IV. <u>In vivo Inhibition of Cryptosporidium parvum</u> Infection

Studies of the inhibitory effect of the polyclonal antibody of the invention on the Cryptosporidium parvum infection in vivo were performed according to the procedure described in Example 17.

HBC 40529 obtained from ImmuCell Corporation (Portland, Maine) was used as a positive control in the animal protection studies described in Example 14 and the HBC Ig of the same lot was also used in the *in vitro* inhibition studies. Assessment of *in vivo* efficacy of HBC was

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performed in newborn, colostrum deprived, Holstein calves challenged with oocysts of Cryptosporidium parvum. The efficacy of the immune colostrum preparation for protecting the treated calves from Cryptosporidium parvum infection was demonstrated in statistically significant differences between treated and control animals in cumulative fecal scores (p< 0.01 by one tailed t test) and dehydration scores (p<0.01 by one tailed t test) (Infect. Immun., 61:4079-4084 (1993)).

As was originally described in the parent application, Ser. No. 07/891,301, incorporated hereby by reference, no dehydration occurred in the treated group whereas all of the calves in the control group showed some signs of dehydration. The occyst output was dramatically reduced in the treated group ($<10^3$ occysts per total fecal output, the limit of detection) when compared to the control group (geometric mean occyst output = 5.62×10^8).

These results clearly show that the immune colostrum treatment effectively reduced the initial colonization by Cryptosporidium parvum parasites and suppressed the intestinal proliferation of the Cryptosporidium parvum parasites which were not initially neutralized.

When similar studies were performed in mice with antibodies to recombinant proteins from the GP900 locus, significant inhibition of infection relative to the positive antibody control (HBC) as well as the negative control was demonstrated with anti-S34, but not with anti-Ag4 or anti- β -galactosidase. Anti-S34 inhibited invasion by 54% percent, whereas the positive control, 1:20 HBC Ig 40529, inhibited by 23% relative to PBS control. This study and the findings confirm that recombinant protein S34 is an effective antigen for the production of passive and active immune products for immunization, prophylaxis and treatment of Cryptosporidium infections.

V. Antibodies and Their Production

Polyclonal or monoclonal antibodies to native or recombinant protein or a glycoprotein of the invention are useful for treatment by providing a protection against Cryptosporidium infections.

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Anti-Cryptosporidium parvum polyclonal antibodies recognizing the cloned polypeptides are preferred over monoclonal antibodies because they recognize multiple epitopes on the target polypeptide.

According to the method of the current invention, large amounts of recombinant polypeptides are easily produced which enable production of a corresponding large quantity of polyclonal antibodies or of the immunogen for active immunization.

The antibodies to recombinant expressed protein can also be produced according to the invention using the standard method available for production of the antibodies to native protein. Some of these method are described in Examples 2, 4, and 5.

VI. <u>Sequences</u>

Six sequences identified as SEQ ID NO 1-6 are disclosed in this invention. These sequences were prepared according to methods described in Examples 10 and 17.

SEQ ID NO: 1 is a DNA sequence of the open reading frame (ORF) of the antigen designated GP900. The sequence comprises 5164 base pairs.

SEQ ID NO: 2 is a DNA sequence of the ORF of the antigen designated GP900 and its 3' flanking region and comprises 5319 base pairs.

SEQ ID NO: 3 is a DNA sequence of the ORF of the antigen designated P68. The sequence comprises 1509 base pairs.

SEQ ID NO: 4 is a DNA sequence of the ORF denoted P68 and its 3' flanking region and comprises 2380 base pairs.

SEQ ID NO: 5 is an ORF sequence of the protein antigen designated as GP900. The GP900 protein contains 1721 amino acids.

SEQ ID NO:6 is an amino acid sequence of the fragment of protein antigen designated as P68. The P68 protein contains 503 amino acids.

VII. Variants and Mutants

Polymorphism found in GP900 is seen in Table 3.

Table 3 represents variant and mutant sequences of

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resulting proteins.

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As seen in Table 3, domains 2 and 4 of the GP900 DNA contain extensive trinucleotide repeats which are expressed as threonine repeat regions. Similar regions occur and have been characterized in the genes responsible for a number of inheritable genetic diseases of man including fragile X syndrome. Insertions and deletions in these regions which are reflected in the translated protein are known to occur. In addition, decreased amount of protein translation has been shown to occur. These protein abnormalities are thought to be related to impaired function of DNA repair enzymes and polymerases in regions of perfect repeats.

A method producing amplified mutants and variants is described in Example 22. Specific variants and mutants vis-a-vis NINC domain 2 of the SEQ ID NO: 5 are shown in Table 3.

TABLE 3 Conservatively Modified Mutants and Variants

of SEO ID NO: 5 Var 2 MGSKVYIPYT KCVGVKH..T TTTTTTTTTT TTTTTTTTT T.....T 5 Var 3 MGSKVYIPYT KCVGVKHTTT TTTTTTTTTT TTTTTTTTTT T....T Var 12 MGSKVYIPYT KCVGVKHTTT TTTTTTTTTT TTTTTTTTTT T.....T 10 Var 2TTT T...... 15 Var 1TTT T...... Var 4 Var 11 TTTTTTTTT TTTTTTTTT TTTTTTTTT TTTTKKPTTT T..... 20 Var 9 TTTTTTTTT AT......TTT T...... 25 Var 11 30 35 40 Var 2 TTTKKPTTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE Var 3 TTTKKPTTTT TTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE 45 Var 12 TTTKKPTTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE Var 1 TTTKKPTTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE Var 4 TTTKEPTTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE Var 11 TTTKKPTTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE 50 Var 9 TTTKKPTTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE Var 10 TTT...TTTT TTTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE TTTTKPTTTT TTTTTTKK PTTTTTATTT TTTSETESVI KPDEWCWLE NINC wherein F=phe W=trp I=ile 55 L=leu R=arq T=thr S=ser P=pro g=gly Y=tyr E=glu H=his Z=OCH or AMB a≖asp Q=qln C=cys A=ala N=asn 60 Z=OPA V=val K=lvs M=met

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NINC sequence seen in the Table 3, corresponds to amino acids 175-423 of the SEQ ID NO: 5. Mutations and variations of the GP900 protein occur within the NINC domain sequence. Corresponding DNA mutations and variations of the NINC bases domain occur in the DNA sequence nt 524-1270 of the SEQ ID NO: 1 and SEQ ID NO: 2.

VIII. <u>Biologically Derived or Recombinant</u> <u>Anti-Cryptosporidium Vaccines</u>

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Vaccine is a biologically derived or recombinantly prepared agent useful for artificially acquired immunization in a host. The current invention describes the production of biologically derived and recombinant vaccines for active immunization of animals and humans against cryptosporidiosis and the preparation of passive immune products for treatment of established infections.

The scope of the invention is, therefore, intended to include biologically derived or recombinantly prepared vaccines based on the antigens of the invention.

A recombinant vaccine is produced by identifying the relevant antigen or antigens of Cryptosporidium, cloning them and expressing them using suitable vectors. approach yields immunogens which are reproducible in sufficiently large quantities to allow preparation of a vaccine for active immunization. Recombinant vaccines are useful for immunization of the potential Cryptosporidium host, such as for inoculation of cows, to produce the host's own antibodies against a Cryptosporidium infection. Additionally, the recombinant vaccines may be used for production of passive immunotherapeutic agents. example, when the cow is inoculated with the vaccine, it begins to produce hyperimmune colostrum. Hyperimmune colostrum from these cows is then purified to yield Ig for passive immunotherapy of immunocompromised primarily AIDS patients.

These vaccines are also useful for widespread use in calves to provide primary protection against Cryptosporidium infection. Providing the herd with anti-Cryptosporidium immunity decreases the risk for waterborne outbreaks of cryptosporidiosis in areas where the watershed

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includes dairy lands. This provides a secondary benefit to humans in those areas. In addition, DNA or RNA may be introduced into a host so that propagation and/or expression of the encoded protein occurs in the host ("a foreign expression system".)

The anti-Cryptosporidium vaccine of the invention contains a Cryptosporidium antigen identified by the invention, modified in such a way that it is incapable of producing the Cryptosporidium symptoms but is capable of eliciting the production of specific protective antibodies against the disease when introduced in the body. A DNA or RNA vaccine for prevention and treatment of infections caused by protozoan Cryptosporidium species (Cryptosporidium) in humans and other mammals was developed by utilizing newly identified and isolated DNA (SEQ ID NOs: 1-4) and amino acid (SEQ ID NOs: 5 and 6) sequences of the Cryptosporidium pathogen designated GP900.

The antigen proteins used for preparation of vaccines correspond to GP900 or to its fragment, or to P68 antigen or its fragment, which are identified by being a target of the polyclonal or monoclonal antibodies of the invention capable of preventing or ameliorating the disease and preventing invasion and/or intracellular development in host cells.

A hybrid vector comprising a DNA segment that encodes the protein antigen able to bind selectively and specifically to anti-Cryptosporidium antibodies operatively coupled to the vector was prepared and expressed. This includes preparation of recombinant vaccines using the viral expression vector outside of the host body and the preparation of DNA vaccines and procaryotic or eukaryotic hosts carrying the hybrid vector which may be introduced into the vertebrate host or a direct introduction of DNA or RNA into host cells generating the host's own expression or translation of DNA or RNA to produce proteins eliciting appropriate antibodies.

Protection from cryptosporidiosis appears to be due to mucosal B cell immunity which, if absent in AIDS patients, is difficult to establish but, if present, may afford

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protection against clinical cryptosporidiosis as AIDS progresses. Thus, the invention describes vaccines able to provide active B cell-immunity against cryptosporidiosis in persons at risk for AIDS or in otherwise immunocompromised patients.

IX. DNA and RNA Vaccines

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Recently, new approaches appeared which utilize so called DNA or RNA vaccines as described in <u>Science</u>. 259:1745 (1993), hereby incorporated by reference.

DNA or RNA vaccines or native immunity are produced according to the methods described <u>Ibid</u>. Briefly, DNA vectors encoding the deactivated anti-Cryptosporidium antigen DNA or RNA are injected, preferably intramuscularly, wherein said antigen is produced and elicits its own immune responses in the form of a specific anti-Cryptosporidium antigen antibody thereby providing its own immunity and/or cell mediated responses.

UTILITY

The current invention provides means for suitable immunoprotection against Cryptosporidium infections or for a therapeutic use of immune agents produced according to the invention.

Immunotherapy and Prophylaxis

The immunotherapy of cryptosporidiosis in humans and animals may be conducted by administration the invention with antibodies of the to patients cryptosporidiosis to effectively reduce their symptomatology.

A method for immunotherapeutic treatment, retardation, or inhibition of Cryptosporidium infection comprises administering to a subject in need of such treatment an amount of an anti-Cryptosporidium polyclonal or monoclonal antibody prepared according to the invention, effective to provide immunity against the invasion of Cryptosporidium or effective to inhibit the existing Cryptosporidium infection.

A method of prophylaxis of Cryptosporidium infection comprises administering to a subject in need of such treatment a vaccine comprising the protein or recombinant

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protein of this invention capable of endogenous development of an inhibitory amount of anti-Cryptosporidium parvum antibodies.

Typical immunization is achieved by inoculation of the animal or human host with the antigen protein combined with equal volume of complete Freunds adjuvant at the beginning of the treatments, with the protein plus equal volume of incomplete Freunds adjuvant at week 2, and with protein combined with equal volume of incomplete Freunds adjuvant at week 4.

For passive immunotherapy Cryptosporidium infected hosts, the polypeptide is first combined with appropriate adjuvants and used for the immunization of cows or other to produce antibodies which animals administered to patients with cryptosporidiosis infection, AIDS patients, and to other particularly to MAb produced in animals, immunocompromised hosts. "humanized" from animal sources, and through chimeric techniques and other derivative techniques may be used for passive immunotherapy.

When in a therapeutic composition, the antigen protein is combined with appropriate adjuvants and used for the immunization of immunocompetent patients who are at risk for cryptosporidiosis either at the time of immunization or in the future. This group includes, but is not restricted to, HIV positive individuals who are still able to respond to vaccination, animal workers, health care workers, day care center children and their caretakers, and children in the developing world.

Formulations suitable for the administration of polypeptides and antibodies such as those described herein are known in the art. Typically, other components stimulatory of immune response may be added as well as fillers, coloring, and the like.

Treatment of Cryptosporidiosis in AIDS Patients and OtherImmunocompromised Subjects

In AIDS patients Cryptosporidium parvum may cause a devastating disease for which there is no treatment. Understanding of the organism and the pathophysiology of

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the disease it produces, and development of treatment, are very important steps in the treatment of Cryptosporidium infection.

Therefore, the protective activity of proteins of the invention was studied in an in vitro Cryptosporidium sporozoite invasion and intracellular development in Madin-Darby Canine Kidney (MDCK) cells. These studies validated the studies performed in the model by correlating inhibition in vitro with protection in an in vivo calf model. The gene for GP900, a sporozoite and merozoite glycoprotein, which is located in micronemes and is accessible to sporozoite surface radioiodination, was identified, cloned and expressed. Similarly, an antigen protein designated as P68 was identified, cloned and expressed. Antibodies raised to fusion proteins expressing portions of the genes (S19, Ag4, S34) inhibited invasion and fusion proteins intracellular development in vitro. Antibodies raised to fusion proteins express portion of the (S19-S34) significantly inhibited intracellular development in vivo in a neonatal mouse model.

GP900 and P68 proteins are therefore the first Cryptosporidium vaccine candidates for preparation of active vaccines and passive immune products for human and animal use in combatting Cryptosporidium infections.

Active vaccines and passive immune products prepared from the protein of the invention are suitable for prevention and therapy of Cryptosporidium parvum infections in AIDS or other patients suffering from cryptosporidiosis.

Currently, no vaccines or pharmaceuticals are available for the prevention or treatment of cryptosporidiosis. Over 95 drugs have been tested in vitro or in vivo, but, none of these drugs has been shown to be effective. One of the therapeutic approaches treatment for of chronic cryptosporidiosis according to the invention is the use of anti-Cryptosporidium hyperimmune antibodies against the protein antigen of the invention which may be given orally to humans to provide a therapeutic benefit.

As the number of AIDS patients increases, the number of cryptosporidiosis cases will also rise resulting in a

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critical need for effective therapy and prophylaxis. The current invention provides an effective treatment and prophylaxis against the cryptosporidiosis infection.

EXAMPLE 1

Cryptosporidium parvum Parasites

This example illustrates the protocol used for isolation of Cryptosporidium parvum parasites.

Occysts of the Iowa and AUCP-1 isolates of Cryptosporidium parvum were passaged through neonatal calves at the Animal Resources Services, University of California, Davis or obtained from a commercial source (Pat Mason) and the occysts were purified and excysted. The detailed protocol is described in <u>Infect. Immun.</u>, 61:4079 (1993).

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Preparation of Murine Anti-Oocyst and Anti-Sporozoite Polyclonal and Monoclonal Antibodies

This example describes the procedure used for preparation of murine anti-occyst and anti-sporozoite polyclonal and monoclonal antibodies.

Polyclonal Antibodies

10 week-old female BALB/c mice were immunized four times intraperitoneally with approximately 5×10^5 sonicated Cryptosporidium parvum oocysts. The polyclonal antibody fraction of the ascites which was shown to react with the Cryptosporidium parvum sporozoite surface, the oocyst surface, and/or with internal antigens of the oocysts, was assessed by an IFA as described in Infect. Immun., 60(12):5132 (1992).

Monoclonal Antibodies

For monoclonal antibody production, mice were immunized intravenously with the supernatant from sonicated Cryptosporidium parvum oocysts three days before fusion as described in J. Immunol., 123:1548 (1979) and J. Parasitol., 68:1029 (1982). Hybridoma supernatants were used as the source of antibodies.

Six sporozoite monoclonal antibodies were obtained. The 10C6, 7B3 and E6 monoclonal antibodies were determined to react with GP900. The supernatants of the corresponding

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hybridoma cultures were used for immunofluorescence assay (IFA) studies and Western blots.

Using the same protocol, infected MDCK cells were used to immunize mice and 3 MAbs to GP900, namely M2, M10, M24 were produced.

EXAMPLE 3

Anti-Cryptosporidium parvum Antibodies Eluted from Western Blot

This example describes the procedures used for preparation of antibodies by elution from Western blot.

For SDS-PAGE, 2x109 oocysts were lysed by 5 cycles of freeze-thawing in 1% Triton Buffer (150 mM Na Cl; 100 mM EDTA; and 1% Triton X-100), in the presence of protease inhibitors: 100 μM E64, 100 μM chymotrypsin, 100 μM pepstatin, and 100 µM leupeptin; and 1.6 mM PMSF, and boiled in sample buffer (SB). Proteins electrophoresed in 5-15% gradient gels described in Nature, 227:680 (1971) and blotted onto nitrocellulose at 0.7 amperes for 8 hours as described in Inf. Immun., 60(12):5132 (1992). Western blots were incubated with HBC Iq (lot # 40529, obtained from ImmuCell Corporation, Portland, Maine) in dilution 1/500 dissolved in 20 ml PBS for 3 hours at 4°C, rinsed 3 times with phosphate buffered saline (PBS), and the antibodies were eluted with 10 ml of glycine buffer (pH 2.6) for 3 minutes, followed by addition of a 1/10 volume of 2M Tris buffer, pH 8 as described in Inf. Imm., 58:2966 (1990). The eluted antibodies were filter sterilized and concentrated to a final volume of 1 ml in a Centriprep 10 concentrator (Amicon, MA).

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Production of Polyclonal Anti-GP900

This example describes the procedure used for preparation of anti-GP900 polyclonal antibodies.

The Triton X-100 (1%) soluble fraction of 2 x 10^8 oocysts was immunoprecipitated with MAb 10C6. A >900 kD MW species was identified in gels stained with Coomassie blue in water and excised. Frozen gel containing 2 x 10^7 oocyst/sporozoites was pulverized and emulsified in 150 PI of PBS and 150 μ l complete Freund's adjuvant (CFA) for

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intraperitoneal (IP) immunization of mice.

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Subsequently, the mice were immunized (IP) three times with the same antigen dissolved in incomplete Freunds adjuvant (ICFA) at approximately 2 week intervals. The anti-GP900 antibody at a dilution of 1:5000 recognized GP900 on Western blots.

EXAMPLE 5

Production of Polyclonal Antibody Against Ag4 and S34 Fusion Proteins

This example describes the procedure used for preparation of the anti-Ag4 and anti-S34 fusion proteins polyclonal antibodies.

Lysogens were produced from the Ag4 and S34 Agt11 clones. Cell lysates and purified protein were made using a protocol and reagents obtained from Promega. Purified fusion protein were emulsified in CFA and injected into rabbits. These injections continued at two week intervals with the substitution of ICFA. Rabbits were sacrificed at the end of 3 months and the antibody was assayed by Western analysis to verify that the antibody recognized a protein >900 kDa.

The β -galactosidase and Ag4- β -galactosidase fusion proteins were purified essentially as described by Promega except that the buffering system used was phosphate buffered saline (PBS) pH 7.4. The purified fusion proteins were then coupled to CNBR sepharose using techniques. The antibodies to Ag4- β -galactosidase were depleted by passaging serum over a CNBR sepharose column coupled to β -galactosidase alone. The flow through fraction was applied to a CNBR sepharose column coupled to the purified Ag4 fusion protein. Antibodies directed against the Ag4 portion of the fusion protein were eluted in 0.1 M glycine at a pH of 2.4 and immediately neutralized in 200 μ l of 2M Tris, pH 7.4. All affinity purified antibodies reacted with the fusion protein and the respective Cryptosporodium protein but not other E. coli proteins.

S34 was subcloned in GST and coupled to a column CNBR sepharose. Antibodies to $S34-\beta$ -galactosidase were passed

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Antibodies directed against the S34 over this column. portion of the fusion protein were eluted in 1M Na thiocyanate and desalted and concentrated.

EXAMPLE 6

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Western Analysis

This example describes the Western analysis method used identify the molecular targets of the protective antibody.

Oocysts (106 lane) were solubilized in 5% β ME (β mercaptoethanol) containing a sample buffer, resolved by SDS-PAGE and subjected to immunoblotting. Proteins were visualized after incubation with primary antibody with 125Ilabeled Protein A followed by autoradiography or with antirabbit IgG conjugated with horseradish peroxidase or followed by calorimetric or phosphatase alkaline chemiluminescent development.

To identify the molecular targets of protective antibody, total Cryptosporidium parvum sporozoite and sporozoite/oocyst proteins were boiled in sample buffer (SB), resolved in 5-15% gradient gels by SDS-PAGE and In blotted with **HBC** Iq. addition, Western sporozoite/cocyst proteins solubilized in Triton-X 100 were immunoprecipitated with HBC Ig at dilutions 1/1,000; 1/5,000; 1/10,000; 1/50,000 and 1/100,000. Cryptosporidium under proteins immunoprecipitated the same parvum conditions but with SHAM-HBC Ig at dilutions 1/1,000 to 1/10,000 were used as controls. Immunoprecipitates were also resolved by SDS-PAGE and Western blotted. blots of HBC Iq immunoprecipitates were developed with HBC Ig (dilution 1:1,000) and SHAM immunoprecipitates were developed with SHAM-HBC Ig (dilution 1:1,000). After incubation with 10 μ Ci [125 I]-protein G for 1 hour at room temperature, blots were dried and exposed for autoradiography.

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EXAMPLE 7

Southern Hybridization

This example describes the Southern hybridization method used for preparation of hybrids of the invention. DNA was purified from 1 x 109 Cryptosporidium parvum 5

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cocysts as described in Example 1. DNA was digested with the restriction enzymes according to procedures provided by the manufacturer Promega. Digested DNAs were subjected to electrophoresis in 0.8% agarose gels in 1X TAE or 0.5X TBE. The gel was blotted to a nylon membrane (Hybond N+, Amersham) per manufacturer's instructions. The probe was labeled with ³²P-ATP and hybridized to the membrane by methods known in the art. Results are seen in Figure 6 where Lanes 1-4 show Iowa isolate DNA and Lane 5 shows AUCP isolate DNA. Lane 1, EcoRI digest; Lane 2, Bgl II digest; Lane 3, Hind III digest; Lanes 4 and 5, Hind I digest.

EXAMPLE 8

Surface Radioiodination and Immunoprecipitation of Cryptosporidium Sporozoite Proteins

This example describes the methods used for surface radio-iodination and immunoprecipitation of Cryptosporidium sporozoite proteins.

Oocysts were bleached, excysted and separated from sporozoites prior to iodination of the sporozoite surface and immunoprecipitation of surface proteins as previously described in <u>Infect. Immun.</u> (1993).

A membrane pellet was prepared by centrifuging 1.1 x 107 sporozoites per ml NETT (0.15 M NaCl, 5mM EDTA, 0.5 M Tris, 0.5% Triton X-100, pH 7.4) at 100,000 x g for 1 hour An aliquot of membrane proteins in 2% SDS 5% pat 40°C. sample buffer was prepared for total sporozoite surface protein analysis. Aliquots of membrane proteins extracted in 2% SDS were diluted with 9 volumes NETT plus 1% high quality bovine serum albumin (BSA) obtained from Sigma; 1 volume 1% Triton X-100; proteinase inhibitors and either MAb 10C6 or anti-GP900 were added for overnight incubation. Protein A Sepharose 4B beads were added to immobilize the immunoprecipitated proteins. Parasite proteins were solubilized in 2% SDS sample buffer containing Bmercaptoethanol. Samples were boiled 5 minutes separated by 5-15% gradient SDS-PAGE.

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EXAMPLE 9

Immunoelectronmicroscopic Localization of GP900 in Cryptosporidium parvum-Infected Rat Intestinal Tissue

This example describes the immunoelectronmicroscopic methods used for localization of GP900 antigen in Cryptosporidium parvum infected rat intestinal tissue.

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Small pieces of terminal ileum were obtained from an immunosuppressed rat experimentally infected with a lamb isolate of the parasite. Tissue samples were fixed with 2% formaldehyde-0.1% glutaraldehyde in PBS for 2 hours at room They were washed in PBS, dehydrated in temperature. ethanol at -20°C, and embedded in LR White obtained from London Resin Co. After polymerization at 37°C for 5 days, thin sections were cut with a diamond knife and collected on nickel grids coated with formvar. They were floated for 30 minutes on 2.5% nonfat dry milk in PBS (PBSM) and then transferred to anti-GP900 mouse ascites obtained described in Example 2 and diluted 1:20 in PBSM for 1 hour at room temperature. After the grids were washed in PBS, they were floated on rabbit anti-mouse immunoglobulin serum obtained from Tago, diluted 1:200 in PBSM, for 1 hour at room temperature, and then transferred for 1 hour to 8 nm protein A-coated beads diluted 1:10 in PBSM.

Thin sections were stained with 3% uranyl acetate in water and observed with a Hitachi H600 electron microscope (EM) Figure 5. EM photographs were also obtained using undiluted MAb IRM hybridoma culture medium and a 1:25 dilution of protein A coated gold beads. Control sections were incubated with unrelated monoclonal and polyclonal antibodies.

EXAMPLE 10

Cloning and Sequencing of a GP900 Locus

This example illustrates the procedure used for cloning and sequencing of a GP900 locus.

The purification and initial characterization of the S34 clone and the description of the restriction fragment genomic expression library from which it was isolated have been described. The Ag4 clone was isolated from the same library as an expression clone which reacted with both the

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anti-GP900 antibody and MAb IRM. The inserts of the S34 and Ag4 clones were subcloned into BlueScript obtained from Stratagene and sequenced in both directions using Sequenase Version 2.0 DNA Sequencing Kit (UBC) or cycle sequencing (New England Biolabs).

DB8, a 3154 bp insert, which contained the sequences of both S34 and Ag4 was identified by a double of screen of the library using these DNA inserts. PCR amplification products for the ends of DB8 and subsequent clones were used to screen the library to identify new clones which extend the sequence 3' and 5'.

EXAMPLE 11

In Vitro Inhibition of Sporozoite Invasion and Intracellular Development

This example describes the methods used for determination of in vitro inhibition of sporozoite invasion/intracellular development.

Oocysts were used to inoculate confluent Madin Darby Canine Kidney (MDCK) cell monolayers for in vitro inhibition assays of sporozoite invasion and intracellular development as previously described in (Inf. Immun., 61:4079 (1993)) with the following modifications. slide wells obtained as tissue culture chamber slides from Nunc Inc., Napersville, Ill, containing 105 MDCK cells were overlaid with 400 l RPMI medium containing 1.5 x 105 oocysts and antibody or colostrum samples to be tested for inhibitory capacity. Each experimental data point was an average of the number of parasite nuclei counted per 200-300 cell nuclei from each of three independently infected Antisera and controls were used after chamber wells. complement inactivation at 55°C for five minutes.

Controls included hyperimmune bovine colostrum 40529 Ig (HBC Ig) raised against *Cryptosporidium* oocysts and sporozoites and SHAM-HBC raised against a herd vaccine at ImmuCell Corp, Portland, Maine.

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EXAMPLE 12

Dose Response Relationship of Affinity Purified Anti-S34 Antibody and Inhibition of Invasion and

Intracellular Development In Vitro

This example describes the method used for determination of the dose-response relationship of polyclonal antibodies in vitro with regard to inhibition of sporozoite invasion and intracellular development.

Affinity purified anti-S34 antibody as described in Example 5 was used to determine the dependence of inhibition of invasion/intracellular development on the quantity of antibody added to the in vitro MDCK assay system as described in Example 11. The antibody, at concentrations of 10, 50, 100 and 500 μ g/ml in RPMI, was incubated with excysted oocysts on MDCK cell monolayers for two hours. The wells were washed out and refilled with RPMI.

Control wells contained equal amounts of oocysts and RPMI alone, S34-GST at 100 nM, anti-oocyst/sporozoite antibody at a 1:40 dilution and HBC Ig 40529 at a 1:40 dilution. As described in Figure 9 invasion/intracellular development was reduced to less than 5% of control in the presence of 500 μ g/ml and less than 20% in the presence of 100 μ g/ml of affinity purified anti-S34 antibody.

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Inhibition of Adhesion by Anti-S34- β -Galactosidase Antibody in the CaCO-2 Adhesion Assay In Vitro

This example describes the method used for determination in vitro of the mechanism by which the polyclonal antibody prevents inhibition of sporozoite invasion and intracellular development.

CaCO-2 cells were grown in monolayers and fixed with paraformaldehyde. Sporozoites were isolated, incubated with 1:50 dilutions of anti- β -galactosidase, anti-S-34- β -galactosidase, anti-Ag4- β -galactosidase and HBC Ig 40529 prepared as described in Example 11. Adhesion was determined using an ELISA assay which had previously been validated by correlation with results determined by

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electronmicrographic assessment of adhesion/inhibition of adhesion.

Anti-S34- β -galactosidase and HBC Ig 40529, the positive control antibody, exhibited an optical density which was 50% of the negative control antibody, anti- β -galactosidase. Anti-Ag4 did not have significant inhibitory activity relative to the control antibody.

EXAMPLE 14

Inhibition of Cryptosporidium Invasion and Intracellular Development in MDCK cells with MAb 10C6

This example describes studies performed to detect inhibition of Cryptosporidium invasion and intracellular development in vitro using monoclonal antibodies.

Cryptosporidium oocysts of the AUCP-1 isolate were excysted and three sporozoite monoclonal antibodies, MAb 10C6, 7B3 and E6, were prepared as described in Example 2. To assess the effect of specific antibodies on sporozoite invasion, MAb 10C6, a monoclonal antibody detecting GP900, was incubated with viable sporozoites for 30 minutes prior to addition to monolayers of MDCK cells.

Sporozoite invasion and intracellular development in MDCK cells was scored at 16 hours after fixation of MDCK cells in formalin and staining with Giemsa. Both invasion and intracellular development were found to be inhibited by > 95% compared to the control antibody. Sequential observation of viable, unfixed Cryptosporidium sporozoites by differential phase contrast microscopy after addition of MAb 10C6 revealed initial reactivity of the MAb with the surface followed by shedding of the sporozoite surface coat and production of a tail-like precipitate. At 30 minutes, shedding was complete or sporozoites were immobile and clumped.

In order to determine whether GP900 was shed by the Cryptosporidium sporozoite in the absence of a specific antibody, living sporozoites were allowed to glide on poly-L-lysine coated microscopic slides. Slides were fixed in formalin and GP900 detected by incubation with MAb 7B3 followed by fluorescein labeled anti-mouse second antibody (Figure 3). MAb 7B3 had previously (data not shown) been

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shown to detect only one protein, GP900, in sporozoites. In Figure 3, the sporozoites were shown to be surrounded by GP900 which was shed posteriorly as the sporozoites glided on the poly-L-lysine coated slides. This reaction occurred in the absence of specific antibody which was added only for detection purposes after fixation of the sporozoites and is analogous to the circumsporozoite deposition and localization of the protein of malaria which contains the binding ligand for binding to the hepatocyte adhesion receptor prior to invasion of the hepatocyte.

EXAMPLE 15

Cloning and sequencing of a portion of the P68 locus

This example describes procedures used for cloning and sequencing of a portion of the P68 locus.

The purification and initial characterization of the S19 clone and the description of the restriction fragment genomic expression library from which it was isolated have been described in Example 10. The S19 clone was subcloned into Bluescript and sequenced. The insert was used as a molecular probe to identify further expression library clones which extended 5' and 3'.

When the sequence data generated was added to that derived from S19, a 2380 bp locus was defined. The locus had 1509 bp of open reading frame which remained open at the 5' end.

Comparison of the deduced protein sequence with the Swiss pro data base indicates substantial homology to mechanoenzymes which may be important in the alteration in shape of the cytoskeleton of the parasite and host during the invasion and establishment of intracellular infection.

EXAMPLE 16

In vitro Inhibition of Sporozoite Invasion/Intracellular Development by Antibody to Recombinant

Proteins from the P68 Locus

35 This example illustrates in vitro inhibition of sporozoite invasion and intracellular development by antibody to recombinant proteins isolated from the P68 locus.

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S19 was subcloned into the pGEX expression vector to yield the expression clone GST-S19, a recombinant protein fused to glutathione-s-transferase. Antibodies were raised to GST-S19 in 2 rabbits (anti-GST/S19#1 and anti-GST/S19#2) the native GST to in 2 rabbits Antibodies anti-GST#2). were assayed in the same inhibition of invasion assay as the GP900 antibodies, however the graph in Figure 12 expresses the results as parasites/MDCK cell nucleus rather than as a percentage of All antibodies were at a dilution of the RPMI control. 1:40 except FCS.

anti-GST/S19 The two antibodies inhibited the Cryptosporidium invasion by 46% and 33% relative to Both were more inhibitory control. than an The anti-oocyst/sporozoite antibody made in rabbits. did inhibit anti-GST antibodies not invasion and intracellular development.

EXAMPLE 17

In Vivo Inhibition of Cryptosporidium Infection in Mice Challenged with Cryptosporidium Occysts with Anti-S34-β-Galactosidase and Anti-S19-β-Galactosidase Polyclonal Antibodies

This example describes the method used for determination of the *in vivo* inhibition of *Cryptosporidium* infection of mice challenged with *Cryptosporidium* oocysts and treated with specific anti-S34- β -galactosidase (Figure 10) and anti-S19- β -galactosidase (Figure 13) polyclonal antibodies.

Anti- β -galactosidase, anti-S34- β -galactosidase, anti-Ag4- β -galactosidase and HBC Ig 40529 were tested for inhibitory activity in a neonatal mouse model of Cryptosporidium infection.

Three experiments were performed and the data pooled. In each experiment 5 neonatal mice per group were infected with Cryptosporidium and were fed either 20 μ l control PBS, 20 μ l of the 3 rabbit antibodies or 20 μ l of a 1:5 dilution of HBC Ig 40529 twice a day. A positive pharmacological control substance, 500 mg/kg/d of paromomycin, in dosage approximately 15 x the dosage given to human AIDS patients

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for cryptosporidiosis, was given to mice in 2 experiments. Infection was scored as the mean number of oocysts shed per day during a 5 day collection period.

Anti-GST antibody (1 rabbit) and anti-GST-S19 antibodies (2 rabbits) were made as described in Example 5, the fusion protein Was glutathione-Stransferase in the pGEX vector. Figure 13 shows the graphical results of a challenge protection experiment in which antibodies were assayed in vivo in groups of 7 CD1 neonatal mice challenged with 104 oocysts orally on day 6. Oocyst output was scored in Sheather's solution and is expressed as 105/ml. Antisera were diluted 1:2 in 50 mM The bars are: 1 is preimmune rabbit sera, 2 is anti-GST antibodies, 3 is anti-S19-GST antibodies from rabbit #1, 4 is anti-S19 antibodies from rabbit #2. Inhibition of occyst excretion relative to control was greater than 45% for the antisera from both rabbits immunized with S19-GST when compared to preimmune sera or antisera to GST alone. When 10 fold more occysts were used in the challenge the inhibitory effect decreased suggesting that the S19 antibodies titrated a particular molecular event.

EXAMPLE 18

Effect of Specific Cryptosporidium Antiqens Mucosal on Lymphocyte Proliferation in Rhesus Macagues Infected with SIV and Cryptosporidium parvum

This example describes the methods used for detection of lymphocyte proliferation specific for S34 and S19 in the mucosa of rhesus macaques infected with SIV and Cryptosporidium parvum.

Rhesus macaques with SIV infection who were used as vaccine controls, that is they were unvaccinated, in an SIV vaccine trial performed at the California Regional Primate Center at Davis, California, were enrolled in a simian AIDS model of a cryptosporidiosis study.

The animals were challenged with whole viable Cryptosporidium oocysts and their clinical state and blood CD4 count were monitored for protracted periods (up to 14 months) before they were sacrificed for evaluation of

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mucosal lymphocyte responses to Cryptosporidium antigens. The animal's entire GI tract was removed and divided into duodenum/jejunum, ileum and colon segments. The tissues were dissected into small fragments, and intraepithelial mononuclear cells (IELs) were released after incubation of the tissue with DTT and EDTA, collected and purified according to the method described in Cell. Immun., 151:379 (1993).

The remaining tissue was digested with Dispase-collagenase obtained from Boeringer-Mannheim to release lamina propria lymphocytes (LPLs). The mononuclear cells released were collected and purified. MNCs were submitted to fluorescent automatic cell sorting with lymphocyte, and macrophage cell marker antibodies. This method shows that the cells collected by the method of Kang are highly enriched for lymphocytes. Results of these studies are seen in Tables 1 and 2 and are described above in section II.

EXAMPLE 19

Agents Suitable for Passive Immunotherapy

This example describes the preparation of passive immunology agents.

The proteins of the invention bind to antibodies which specifically bind to epitopes of Cryptosporidium parvum. These Cryptosporidium parvum epitopes are also recognized by B and T cells. The proteins mentioned above are in amounts reinserting the produced large by Cryptosporidium parvum DNA from the different clones described in Section I, above, into an expression vector such as pGEX, pET-9d, or baculovirus. The thus constructed hybrid vector is used to transform or transfect a host. The host cells carrying the hybrid vector are then grown in a nutrient medium to allow the production of the gene product.

Vectors pGEX (Pharmacia), disclosed in <u>Gene</u>, 67:31 (1988) or pET-9d (Novagen)/pRSET T7 (Invitrogen) utilize the T7 RNA polymerase and the T7 promoter according to <u>Meth. Enzymol.</u>, 185:60 (1990) and hosts derived from *E. coli*. Following protein expression, the vector sequences

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are easily eliminated so that the subsequent immunogenic protein contains only Cryptosporidium sequences. These expression systems are commercially available and their use is standard in the art.

Recombinant baculovirus is a simple vehicle for the expression of large quantities of protein from eukaryotic or prokaryotic gene origin. The genes are expressed under the control of the Autographa californica multiple nuclear polyhidrosis virus (AcMNPV) polyhedral promoter contained in transfer vectors used to infect Spodoptera frugiperda (Sf9 or Sf21) insect cells.

A number of transfer vectors is available for the production of protein from both full length and partial cDNA and genomic clones. Fused or non-fused protein products, depending on the vector used, can produce up to 50% of the total protein in infected cells. The thus obtained recombinant proteins are frequently functionally similar immunologically and to corresponding endogenous proteins. Proteins with signal peptides may be secreted into the media while those without secretion signals will aggregate in the cells or be localized at the membrane. Baculovirus expression systems are commercially available from Invitrogen.

Thus obtained polypeptide is purified by methods known in the art, and the degree of purification varies with the use of the polypeptide. For use in eliciting polyclonal antibodies, the degree of purity may not need to be high. However, as in some cases impurities may cause adverse reactions, certain degree of purity is preferred and required.

EXAMPLE 20

Agents Suitable for Active Immunotherapy

This example illustrates the agents suitable for active immunotherapy.

Peptides, polypeptides, glycopeptides or proteins comprising epitopes of Cryptosporidium parvum recognized by B and/or T cells are produced in large amounts by recloning, as described in Example 19, above. The polypeptide thus obtained are purified as described above.

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The degree of purification varies with the use of the polypeptides. For use in eliciting polyclonal antibodies, the degree of purity may be lower than for other applications. For the preparation of a pharmaceutical composition, however, the degree of purity must be high, as is known in the art.

When in a therapeutic composition, the polypeptide is combined with appropriate adjuvants and used for the immunization of immunocompetent patients who are at risk for cryptosporidiosis either at the time of immunization or in the future.

This group includes, but is not restricted to, HIV positive individuals who are still able to respond to vaccination, animal workers, health care workers, day care center children and their caretakers, and children in the developing world.

In alternative, the peptides, polypeptides, glycopeptides or proteins of the invention are prepared synthetically using methods known in the art. General methods for synthesizing peptides, polypeptides, glycopeptides or proteins are described in U.S. Patent 5,527, 882, incorporated hereby by reference.

EXAMPLE 21

Preparation of Anti-Cryptosporidium parvum Vaccine

25 This example illustrates procedure for anti-Cryptosporidium parvum vaccine.

Vaccine use of recombinant Cryptosporidium antigens.

- (1) Antigens: Preferably 10-200 μ g of recombinant antigen of the invention, either alone or in combination.
- 30 (2) Adjuvant: Any one of a number of adjuvants designed to either:
 - (a) stimulate mucosal immunity;
 - (b) target mucosal lymphoid tissue.

Some examples are: liposomes, saponins, lectins, cholera toxin B subunit, E. coli labile toxin (LT) B subunit, pluronic block copolymers, hydroxyapatite, plant glucans, acetyl mannan (from Aloe Vera), aluminum hydroxide.

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- (3) Route of administration: Since the vaccine must stimulate mucosal immunity, it preferably is delivered to a mucosal site. Feasible routes of administration include: oral, nasal, rectal, and vaginal. However, boosting may occur via another route, for example, intramuscular or subcutaneous, or may involve the use of other methods such as foreign DNA which replicates in vivo and dictates protein expression in the host.
- (4) Volume: The volume of the vaccine, while not particularly important, should be in the range that would permit ease of use. Preferred range would be 0.5 ml-2.5 ml or so (including adjuvant).
 - (5) <u>Boost schedule</u>. For non-immunocompromised individuals, the standardly used booster schedule is used.

For immunocompromised individuals, a more aggressive boosting schedule is used. The vaccine is administered to high risk patients initially when their immune status is reasonably good (i.e., CD4 count of >500).

The initial booster is given 1 month after the primary immunization, and again every 3-4 months during progression of the immunodeficient state.

EXAMPLE 22

Polymorphisms in GP900

This example illustrates the method used to prepare mutant and variant products.

Genomic DNA from the Iowa and several other strains was subjected to PCR amplification using primers which were situated outside of domain 2, in the distal region of domain 1 and the proximal region of domain 3. Several prominent bands of different sizes were observed when the PCR products were visualized by ethidium staining of a gel in all of these strains. As a control for tac polymerase, DB8 DNA was also amplified by the polymerase chain reaction. Only a 700 bp amplification product was detected indicating that the multiple bands were a product of amplification of sequences present in the genomic DNA, and were not an artifact of the PCR process.

Two of the amplification products were cloned into sequencing vectors and 4 clones from each of the products

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were sequenced to determine their relationship to the NINC domain 2 sequence (Table 1). All 8 sequences had an open reading frame indicating that they were portions of DNA which could be the blueprint for a GP900 protein. sequences appeared to have in-frame (multiples of 3) DNA deletions with respect to the NINC sequence. All 8 coded for a domain 2 which had conservation of the threonine rich regions, but all 8 differed from each other. The DNA data indicate that mutation in domain 8 is common. conservation of threonines and the in-frame nature of all 8 clones indicate that there are selection pressures acting at the level of the protein (presumably production of an attachment protein which will mediate attachment and allow invasion and propagation) which determine which genotypic variants are maintained in a strain. show that there are mutants of GP900 which are maintained in an isolate's gene pool, presumably as variant alleles at a single locus in haploid stages of the organism.

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SEQUENCE LISTING

(1) GENERAL INFORMATION: THE REGENTS OF THE (i) APPLICANT: UNIVERSITY OF CALIFORNIA VACCINES, ANTIBODIES, (ii) TITLE OF INVENTION: PROTEINS, GLYCOPROTEINS, DNAS AND RNAS FOR PROPHYLAXIS AND TREATMENT OF Cryptosporidium parvum INFECTIONS (iii) NUMBER OF SEQUENCES: (iv) CORRESPONDENCE ADDRESS: ROBBINS, BERLINER & CARSON, (A) ADDRESSEE: L.L.P. 201 N. Figueroa Street (B) STREET: 5th Floor Los Angeles (C) CITY: California (D) STATE: United States of America (E) COUNTRY: 90012-2628 (F) ZIP: (v) COMPUTER READABLE FORM: Diskette - 3.5 inch, 1.44 Kb (A) MEDIUM TYPE: storage PC (B) COMPUTER: DOS (C) OPERATING SYSTEM: Wordperfect 5.1 (D) SOFTWARE: (vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: (C) CLASSIFICATION: (vii) PRIOR APPLICATION DATA: 08/415,751 (A) APPLICATION NUMBER: April 3, 1995 (B) FILING DATE: (viii) ATTORNEY/AGENT INFORMATION: (B) REGISTRATION NUMBER: 20.121 (C) REFERENCE/DOCKET NUMBER: 5555-449C1 (ix) TELECOMMUNICATION INFORMATION:

(213) 977-1001

(213) 977-1003

(2) INFORMATION FOR SEQ ID NO. 1:

(A) TELEPHONE:

(B) TELEFAX:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 5163 base pairs

(B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Cryptosporidium parvum

(xi) SEQUENCE DESCRIPTION: SEQ ID NO. 1:

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	ACTACTGGAT					120
	CTGAAGGATC					180
	GGTTATAGCA					240
	GATCCATACT					300
	GGTAAAACTA					360
	GAGCCTAGTG					420
	CCTAGTGCAA					480
	GAGAACTCTT					540
	ACTAAATGTG					600
	ACGACAACAA					660
CACTACTACT	ACGACAACAA	CAACAACAAC	AACAACAACA	ACAACAACAA	CAACAACAAC	720
AACAACAACA	ACAACGACTA	CTACTACTAC	TACTACTACT	ACTACTACTA	CTACTACTAC	780
AACCACAACA	ACTACAACCA	CAACTACAAC	CACAACAACT	ACAACCACAA	CAACAACAAC	840
CACAACAACT	ACAACCACAA	CTACAACCAC	AACAACTACA	ACCACAACAA	CCACAACCAC	900
	ACCAAGAAAC					960
	ACCACCACAA					020
	ACTACTACCA					1080
	ACTACAACAA					1140
	ACAACAACTA					1200
	ACAACTACTA					1260
					GTGTTATCGG	1320
					ATGACACGCA	1380
					GATGCAGAAA	1440
	AAACTCGAGT					1500
					TTCACGTAAG	1560
					CTGAGTTATT	1620
					TGGATATCAA	1680
	ATTTCGTAAC					1740
					TACTACCACT	1800
					AACTCAAAAA	1860
					AACAACAACA	1920
					AACAACAACA	1980
					GACACCATTA	2040
					GGATAAGTAC	2100
					TGAACCAATT	2160
					TCAAACTCAC	2220
					AAAATCAGGT	2280
					ATATETTECT	2340
					TACACTCACT	2400
GGTTACCCAT	TUGATECAGT	CAGITIGATI	CCGTTCAATC	CAGAAACTG	TGAATTGTTT	2460

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GATCCAATAT	CAGATGAGAT	AATGAATGGA	ACAATTGCAG	GTATTGTTTC	AGGAATTTCT		2520
GCAAGTGAGT	CATTATTATC	TCAGAAATCA	GCTCTAATCG	ACCCAGCAAC	AAATATGGTT		2580
GTTGGAGAAT	TTGGTGGATT	GTTGAACCCA	GCAACAGGAG	TGATGATTCC	AGGTTTTTTA	;	2640
GGTCCATCAG	AGCAAACTCA	ATTCTCCCCT	GAGATTGAAG	ATGGTGGTAT	TATTCCTCCA		2700
GAAGTAGCAG	CAGCAAATGC	TGATAAATTC	AAGTTATCTA	TTCCTCCAAG	CGTACCAGAA		2760
TCAATTCCAG	AAAAGGATCA	GAAGATTGAT	TCTATTTCTG	AATTGATGTA	TGATATTGAG		2820
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CCAACCACAG	GTCTGCCTTT	CAATCCACCA	ACTGGTCATT	TGATTAACCC	AACAAATAAT	:	3000
• • •	ATTETTCATT					;	3060
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TCAGATATTC	CATTTAACTC	AACTACAGGT	GAATTAGTTG	ATCCATCAAC	AGGAAAGCCA		3180
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	CAGTCAATGG						3420
GATAAGGGTA	AGGATGGATT	AATTGTTCCA	CCAACTAATT	CTATCAATAA	AGATCCAGTA	:	3480
ACAAATACTC	AGTACAGTAA	TACTACTGGT	AACATTATTA	ACCCAGAAAC	AGGAAAAGTT	:	3540
ATTCCAGGTT	CACTTCCAGG	CTCTCTCAAC	TATCCATCAT.	TCAATACTCC	ACAACAAACT		3600
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	AAGTATTGAA						3780
	TTCCAAGAGA						3840
	CAAATAAAAA						3900
	AATCACATGA						3960
	CATTCGATGA						4020
	CACATTCTGG						4080
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TCTGCAGCTT	CAGTAATTCA	CACTGCTCTT	GGAACACCAA	CTCAAACTGA	TCCAACAACA	43	320
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GATCCTCAGA	CTGGAGAGCA	AATCAAGGGT	TCTGTTCCTT	ATGTTTCATT	GTACGTTAAG	44	440
GAAAAGAATA	TTGTAACAGA	AGCTGCTTAT	GGTCTACCAG	TTGATCCAAA	GACTGGTTTC	45	500
CCAATTGATC	CAATTAGTTA	CCTCCCGTTT	GCTAAGAATG	GCGAACTAAT	TGATCCTATC	45	560
TCTGGTAAAT	ATTTCAGTGG	TTCAATTGCT	GGATTCATTT	CTGGTAAAGC	TGGTTCACAA	46	620
TCTAAATCAT	CTGATGAATC	AGGTAATCCA	ATTGATCCAT	CAACAAATAT	GCCTTACGAT	46	680
CCAAAAGGCG	GCAAATTAAT	TGATCCAGAA	TCTGGCATTG	CTATTGATAA	TTCTGTTTCA	47	740
GGTGTGTTTG	CAACTGTACC	TGGTACTGCT	GCACCGAAAA	AGGGTGGTGT	CATTCCGGAG	41	B00
TCAGTTGCAG	CTGAGGCAGC	AAAGAAATAC	TTTGCAGCCA	ATGTTGAGGG	AGAGGGAGAA	48	B60
GGAGAAGAAG	TTCCACCACC	GCCAGAATCA	TCTAGTAACA	TTGCAATCCA	AGCTGCTGGT	49	920
GGTGCTTCTG	CTGCTGTAGG	TCTCGTAGCT	GCTGTTGGTG	CATGGTATGC	AAGCAGAAAC	49	980
AGACAGGAAG	GAGAAGATGA	TGATGACTAT	CAGATGGATT	TGAAGCAGAA	TATGAAGAAG	50	040
AAGAGGAAGA	AGAGGGTGAT	GAAGCAGCAA	ATGAAACTGT	TGTTACAATT	GAGCGTGATT	5'	100
CATCATTCTG	GAACGAATCT	TAAACGTAGA	AAAGATTTTT	CCAATTCAAA	AAAATTTCGA	5·	160
ATA						5'	163

(3) INFORMATION FOR SEQ ID NO. 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 5318 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: doubte
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (vi) ORIGINAL SOURCE:
 (A) ORGANISM: Cryptosporidium parvum
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO. 2:

AATITIGGAA GGTICAATIG CAGGTATTAG AAGCGAATCT TGCATTGTAT CTGAACTGAA	60
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AAGTGGTGAA CGTAAGGATC CAAACAAACA AGCTACTATT TCTGGTTCAA GATCTTGTGG	180
ATGGAAACAA GGTTATAGCA TTGATTCATC CACCGGGTTT AGAGTTGATT CTATCACTGG	240
TETECCAACT GATCCATACT CTAATTGTCC ATTCAACCCT GTCACTGGAA ATTTAGTCAG	300
TAGGTECACT GGTAAAACTA TTCCAAACAC TTATGCAGGT GTTTATCGTT CTAATGAGAC	360
TAAGACCACT GAGCCTAGTG CAAACACTTA TGCAGGTGTT TATCGTTCTA ATGAGACTAA	420
GACCACTGAG CCTAGTGCAA ACACCAACTT CTTGTTGGTA GATCCTAAGA TTAATGCTCC	480
TIGTAATICT GAGAACTCTT TIGAACAAGG TCAAATATIT GATATGGGCA GTAAGGTATA	540
CATTCCATAC ACTAMATGTG TTGGAGTGAA ACACACAACA ACAACAACAA CAACTACTAC	600
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ACTAAGGCAA CAACAACCAC AACAACTCTT AATCCAATCA TTACAACAAC AACTCAAAAA	1860
CCAACAACAA CAACAACAAC AAAGGTTCCA GGTAAGCCAC CAATAGCCAC AACAACAACA	1920
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GTGCCAACGA CAACTACTAC TACCAAGAGA GACGAAATGA CAACAACAAC GACACCATTA	2040
CCTGATATCG GTGACATTGA AATTACACCA ATCCCAATTG AAAAGATGTT GGATAAGTAC	2100
ACAAGAATGA TITATGACTA TAACAGTGGT TTATTATTAG ACTCTAATGA TGAACCAATT	2160
CCAGGTTCTC AAGCAGGACA AATAGCTGAT ACAAGCAATT TATTCCCAGT TCAAACTCAC	2220
AAGAGTACTG GTTTACCAAT TGATCCAATG GTTGGTCTTC CATTTGATCC AAAATCAGGT	2280
AATTTAGTAC ATCCATATAC CAATCAAACA ATGTCTGGTT TATCGGTATC ATATCTTGCT	2340

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	TGGATCCAGT					•	2460			
	CAGATGAGAT						2520			
	CATTATTATC						25 8 0			
GTTGGAGAAT	TTGGTGGATT	CTTGAACCCA	GCAACAGGAG	TGATGATTCC	ACCTTTTTA	· · · · · · · · · · · · · · · · · · ·	2640			
	AGCAAACTCA						2 70 0			
	CAGCAAATGC						2760			
	AAAAGGATCA					· · · · · · · · · · · · · · · · · · ·	2820			
	TTATTGGTCA						2880			
	TAATGAAGAC						2940			
	GTCTGCCTTT						3000			
AATACCATGG	ATTCTTCATT	TGCTGGTGCA	TACAAATATG	CAGTTTCAAA	TGGTATTAAG		3060			
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	CATTTAACTC						3180			
ATTAACAATT	CTACTGCTGG	TATTGTTAGT	GGAAAACCTG	GCTTACCACC	TATTGAAGAT		3240			
	ATTTGTTTGA						3300			
	AAACCAACAG						3360			
	CAGTCAATGG						3420			
	AGGATGGATT						3480			
	AGTACAGTAA						3540			
ATTCCAGGTT	CACTTCCAGG	CTCTCTCAAC	TATCCATCAT	TCAATACTCC	ACAACAAACT		3600			
GATGAGATTA	CAGGAAAGCC	AGTTGATACT	GTTACTGGTT	TGCCATATGA	TCCATCTACA		3660			
GGTGAAATTA	TCGATCCTGC	AACTAAATTA	CCAATTCCAG	GATCAGTTGC	AGGTGATGAA		3720			
ATCCTCACTG	AAGTATTGAA	CATTACAACA	GATGAAGTAA	CAGGTTTGCC	AATTGATCTT		3780			
	TTCCAAGAGA						3840			
GTTGATCCAT	CAAATAAAAA	ACCAATTCCA	GGTTCACATT	CCGGATTTAT	TAATGGTACA	•	3900			
	AATCACATGA						3960			
GGTTTGCACC	CATTCGATGA	AGATTCAGGT	AGTTTAATTA	ACCEAGAGAC	TGGAGATAAA		4020			
	CACATTCTGG						4080			
	TGACACCTGA						4140			
	TTTCACCAAG						4200			
	AGATTTCTGG						4260			
	CAGTAATTCA						4320			
	CTGATCCATC						4380			
	CTGGAGAGCA						4440			
	TTGTAACAGA						4500			
	CAATTAGTTA						4560			
	ATTTCAGTGG						4620			
	CTGATGAATC						4680			
	GCAAATTAAT						4740			
					CATTCCGGAG		4800			
					AGAGGGAGAA AGCTGCTGGT		4860			
					AAGCAGAAAC		4920			
					TATGAAGAAG		4980			
					GAGCGTGATT		5040 5100			
					AAAATTTCGA					
					TTCCTATTGA		5160			
					AAAATTCAAC		5220			
	ATCTATCAAA			TAMINATGAC	MAMATICANU		5280			
ANINI GAIGA	AIGIAIGAAA	aca i i caa	I GUNUNAN				5318			
		-								
(4) INFORM	(4) INFORMATION FOR SEQ ID NO. 3:									
(i) SEQUENCE	CHARACTERIS	STICS:							

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1509 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

- (vi) ORIGINAL SOURCE:
 (A) ORGANISM: Cryptosporidium parvum
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO. 3:

AGTAAGGGTC AATTATTYAA CCCAGTAAGT AAGTTGTGTG TACGACTTAA AGACAATGTT	60
GTAGGTGGAG GAGCTCTGGT TTTGGATGAT TGTCGTAAAG CTAGTGATGG AAGTGGATTA	120
TTCGAATTAA TGCCAAACAA TCAGCTCAGA TTAGCTAGAG GTGGAAATCT ATGCTTAACA	180
AGTCCAGGAG ATAAGCCAGG AGTCGCGAAT GTTGCATTAA ACTCAGCAGC CAGTTCCACA	240
AGTGTGGTTA GAACAGGTAT TGAGAATGGT CCAGCAATGG CTGTTGATGG AAAGGATACA	300
TCATATTGGT TGTCAGATTC TTCAACTCCT GGTAAAGATT CTGCAAATGT TAACTTTTTG	360
AGTGATACCG GTTCAGTTAC AAAACTTAAA GATATCTTTA TTGAGTGGAA ATATCCTGCC	420

ATTGACTTTA	ATATTGATTT	AAGTGAAAAT	GGAAAGGAAT	ATCAAACCCA	AGTTTCTGTG	486
AATAATAATG	GATTAATGTC	AACCACTTAT	TCATTAGAAG	GAAAGAAAGC	AAGATATGTC	54
AAAATTCAAA	TGACAATTCC	AAGCCAAGAT	GAGACAGGGA	AATATGTGTA	TGGTATCAAA	600
CAGGTGAGAA	TATTCAGTAA	TACTATGAGA	AGTACTGTTG	AGGATTGTAG	TAGTGTTAAA	666
CAACATAATG	ATGGTAGAGA	CAAAATATTC	CCACTCCCAT	ATAATGGTGA	TAATTTTGCA	72
CCCGGATTAT	TGTTAAAGGC	TCACGGAATT	AGTGTAAAGA	ATAGATTAAA	TGAATTACAA	78
GAGCTTTCTG	GTAAGGTAAC	TTCAATATTA	CCAAATTTGG	ATGCATGTAG	AAAGACTTCT	841
GATGGAAGAG	ATAACACATT	AAAGATGCAG	GCAACCAAAT	TAGGATTTTT	GTCAGAAAAA	900
TTGGAGAAAT	TAACTTCCGA	CTATAATCTC	GAGTATAAGT	TTACGAAGCC	AGCTTTAGGA	961
GGTTCCGAGT	TATATCCAGG	GGAAGATTGT	GTTGCTATTA	AGAATGATAA	GACTCAGGAA	102
GCCATTAGTG	GTTTTTATTA	TGTTAGACCA	TTCTGTTCAA	CCAAACCATT	GAGAGTTTAC	108
TGTGATATGA	ACACTGGAAA	TACAATCTAT	CCAATGGAAA	TGAGTGTTCA	TTCTTCCAGA	1140
GCAGCTTCTT	CAGCATGTGC	AACTGTTGGA	TTAAAACCAT	TATTGTTAAG	GGACAAAAAG	120
GAATCTGTTG	TAGGTATTAA	GAAGATGTTG	AATATGATGA	ATATTAATGA	TAATAGAAGA	126
GTTATTCCTT	TGACTCACGA	CTTTGGTTGT	GATAATCCTA	AAGGATGCAA	TTCACAATTT	132
ACACAGTTAG	GCAGTGGTGT	TGAAGAATTT	GTTGCTGCAT	CTCCTCAGGC	AGCAGCTTCA	138
AACTCTACAT	CTGGAGCACT	TCCAGAACTG	GTTCTTTGCA	GTACAAATAC	CAATTTGAAG	144
	ATGCAATTTC					150
CATTTGGAT						150
						130

(5) INFORMATION FOR SEQ ID NO. 4:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2380 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA (genomic)

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Cryptosporidium parvum

(xi) SEQUENCE DESCRIPTION: SEQ 1D NO. 4:

AGTAAGGGTC	AATTATTAA	CCCAGTAAGT	AAGTTGTGTG	TACGACTTAA	AGACAATGTT		60
GTAGGTGGAG	GAGCTCTGGT	TTTGGATGAT	TGTCGTAAAG	CTAGTGATGG	AAGTGGATTA		120
TTCGAATTAA	TGCCAAACAA	TCAGCTCAGA	TTAGCTAGAG	GTGGAAATCT	ATGCTTAACA		180
AGTCCAGGAG	ATAAGCCAGG	AGTCGCGAAT	GTTGCATTAA	ACTCAGCAGC	CAGTTCCACA		240
AGTGTGGTTA	GAACAGGTAT	TGAGAATGGT	CCAGCAATGG	CTGTTGATGG	AAAGGATACA		300
TCATATTGGT	TGTCAGATTC	TTCAACTCCT	GGTAAAGATT	CTGCAAATGT	TAACTTTTTG		360
AGTGATACCG	GTTCAGTTAC	AAAACTTAAA	GATATCTTTA	TTGAGTGGAA	ATATCCTGCC		420
ATTGACTTTA	ATATTGATTT	AAGTGAAAAT	GGAAAGGAAT	ATCAAACCCA	AGTITCTGTG		480
AATAATAATG	GATTAATGTC	AACCACTTAT	TCATTAGAAG	GAAAGAAAGC	AAGATATGTC		540
AAAATTCAAA	TGACAATTCC	AAGCCAAGAT	GAGACAGGGA	AATATGTGTA	TGGTATCAAA		600
CAGGTGAGAA	TATTCAGTAA	TACTATGAGA	AGTACTGTTG	AGGATTGTAG	TAGTGTTAAA		660
CAACATAATG	ATGGTAGAGA	CAAAATATTC	CCACTCCCAT	ATAATGGTGA	TAATTTTGCA		720
CCCGGATTAT	TGTTAAAGGC	TCACGGAATT	AGTGTAAAGA	ATAGATTAAA	TGAATTACAA		780
GAGCTTTCTG	GTAAGGTAAC	TTCAATATTA	CCAAATTTGG	ATGCATGTAG	AAAGACTTCT		840
GATGGAAGAG	ATAACACATT	AAAGATGCAG	GCAACCAAAT	TAGGATTTTT	GTCAGAAAAA		900
TTGGAGAAAT	TAACTTCCGA	CTATAATCTC	GAGTATAAGT	TTACGAAGCC	AGCTTTAGGA		960
GGTTCCGAGT	TATATCCAGG	GGAAGATTGT	GTTGCTATTA	AGAATGATAA	GACTCAGGAA	•	1020
GCCATTAGTG	GITTITATTA	TGTTAGACCA	TTCTGTTCAA	CCAAACCATT	GAGAGTTTAC	•	1080
TGTGATATGA	ACACTGGAAA	TACAATCTAT	CCAATGGAAA	TGAGTGTTCA	TTCTTCCAGA	•	1140
GCAGCTTCTT	CAGCATGTGC	AACTGTTGGA	TTAAAACCAT	TATTGTTAAG	GGACAAAAAG	•	1200
GAATCTGTTG	TAGGTATTAA	GAAGATGTTG	AATATGATGA	ATATTAATGA	TAATAGAAGA	•	1260
GTTATTCCTT	TGACTCACGA	CTTTGGTTGT	GATAATCCTA	AAGGATGCAA	TTCACAATTT	•	1320
ACACAGTTAG	GCAGTGGTGT	TGAAGAATTT	GTTGCTGCAT	CTCCTCAGGC	AGCAGCTTCA	•	1380
AACTCTACAT	CTGGAGCACT	TCCAGAACTG	GTTCTTTGCA	GTACAAATAC	CAATTTGAAG	•	1440
CATGAAAGCA	ATGCAATTTC	CTTGTCTTGT	GAAAGCAGAT	TCTCTGATAT	GAAGGTATTT	•	1500
CATTTGGATT	AGTAACCTGA	ATTAAATGAT	GTAGAAGAGA	TCTAATAGCT	TTAGTATGTT	•	1560
GCAAAAATTC	GTTAGAAAGT	TCAAGGAACT	CAAGCTTAAA	CTTTCTTGTT	TICTITCTCC	•	1620
ATGATTTTT	CTTGTTATAC	TTCTCTGCAA	CCCTAAGTGT	TTCTTGCCCA	AAATTGATTA		1680
ATTCTCTGAT	CTTGGGCCTA	TATTTGGTGA	ACTCATTAAT	AAATGTTTCA	GTAACGACAC		1740
TATTTGGGTT	TTCTGTTAAA	TCTAGTCCAA	AGTTAACAGT	TTTGATCTTC	TTTGCCTTAT		1800
TATAACAAAC	ATTAATCTCC	TTTTCTTGTT	CAGGAGATAG	TTCAACGTCT	TTTGAGTATG		1860
AAATACTTGC	TTCTTTATTA	TAATTCGGAT	ATTCGCTTTG	ATCAGTTTGG	CCTTCTGATC		1920
CGTCAATTTG	TGATCTACGT	TCTATAATAG	CTTCTGGAAT	ATCTGTAGCA	GGAGCATCTT		1980
GCGGCTTTAA	TCCAACAGGA	AGCTCCTTTA	CTGTATTAAT	ATAGGAGAAC	GGCATATTTC		2040
CAAAGCCATT	AGTATTTAAT	TCGCCATTAT	TCTGAGTACC	CTTGGGAGAA	TTATCGCTAA		2100
TAGAATCCCC	TTCAGCCTTT	TGAAGGTTGA	AATTCAGCTC	AGTTTCATCO	ATACTACTTG		2160
GATCAGAATC	TTCAAATCCA	GTAGCTAGT	TTCTAGAGA	AGACTCAGCA	TCACTATATC		2220
CTTCATTAGT	TGTTTCCTCA	GATTCTAGA	A TCTTCTCCG1	GATACTITCA	GTAGGATTTG		2280

ATGCGCGTAA ATACAGGGCT TICCTGCTTG TIGAAATGGC CAGTIICTGT AATIIGAGTT TTTTCCTCAC TITCAGACTG TICTGGATAA TCCGGAATTT

2340 2380

- (6) INFORMATION FOR SEQ ID NO. 5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1721 amino acids
 - (B) TYPE: amino acid (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Cryptosporidium parvum
 - (ix) FEATURE
 - (A) NAME/KEY: antigen
 - (B) IDENTIFICATION METHOD: deduced
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO. 5:

ile leu glu gly ser ile ala gly ile arg ser glu ser cys ile vat 1 5 10 15 glu leu asn phe thr ser thr thr gly phe thr thr asp thr ser 20 25 30 met asn trp pro val ser ile thr ser gly glu arg lys asp pro asn 35 40 45 gln ala thr ile ser gly ser arg ser cys gly trp lys gln gly 50 55 60 tyr ser ile asp ser ser thr gly phe arg val asp ser ile thr gly 65 70 75 80 leu pro thr asp pro tyr ser asm cys pro phe asm pro val thr gly
85 90 95 asn leu val ser arg ser thr gly lys thrile pro asn thr tyr ala 100 105 110 gly val tyr arg ser asn glu thr lys thr thr glu pro ser ala asn 115 120 125

thr tyr ala gly val tyr arg ser asn glu thr lys thr thr glu pro 130 135 140 ser ala asn thr asn phe leu leu val asp pro lys ile asn ala pro 145 150 155 160 cys asn ser glu asn ser phe glu gln gly gln ile phe asp met gly 165 170 175 ser lys val tyr ile pro tyr thr lys cys val gly val lys his thr 180 185 190 thr thr thr thr thr thr thr thr thr lys lys pro thr thr thr thr thr 585 390 395 400 ala thr thr thr thr thr ser glu thr glu ser val ile lys pro
405 410 415

asp glu trp cys trp leu glu lys asn gly glu cys glu ala lys gly
420 425 430 ata thr tyr val gly vat ite gly lys asp gly arg ite glu asn gly
435 440 445 met ala phe thr met ile pro asn asp asp thr his val arg phe arg 450 460 phe lys val lys asp val gly asn thr ile ser val arg cys arg lys 465 470 475 480 gly ala gly lys leu glu phe pro asp arg ser leu asp phe thr ile 485 490 495 pro pro val ala gly his asn ser cys ser ile ile val gly val ser 500 505 510 gly asp gly lys ile his val ser pro tyr gly ser lys asp val ser 515 520 525 leu ile ser ala pro ile gln pro ser glu leu phe asn glu val tyr
530 535 540 cys asp thr cys thr ala lys tyr gly ala ile his ser gly tyr gln 545 550 555 560 thr ser ala asp phe val thr thr thr thr ala lys pro thr thr thr 565 570 575 thr thr gly ala pro gly gln pro thr thr thr thr thr gly ser pro 580 585 590 ser lys pro thr thr thr thr thr thr lys ala thr thr thr thr thr 595 600 605 thr leu asn pro ile ile thr thr thr thr gln lys pro thr thr thr 610 615 620 thr thr thr lys val pro gly lys pro pro ile ala thr thr thr thr 625 630 635 640 thr leu lys pro ile val thr thr thr thr thr lys ala thr thr thr 645 650 655 thr thr thr val pro thr thr thr thr thr thr lys arg asp glu
660 665 670 met thr thr thr thr pro leu pro asp ile gly asp ile glu ile 675 680 685 thr pro ile pro ile glu lys met leu asp lys tyr thr arg met ile 690 695 700 tyr asp tyr asn ser gly leu leu leu asp ser asn asp glu pro ile 705 710 715 720 pro gly ser gln ala gly gln ile ala asp thr ser asn leu phe pro 725 730 745 val gln thr his lys ser thr gly leu pro ile asp pro met val gly 740 750 leu pro phe asp pro lys ser gly asn leu vai his pro tyr thr asn 755 760 765 gin thr met ser gly leu ser val ser tyr leu ala ala lys asn leu 770 780 thr val asp thr asp glu thr tyr gly leu pro ile asp thr leu thr 785 790 795 800 gly tyr pro leu asp pro val ser leu ile pro phe asn pro glu thr 805 810 815 gly glu leu phe asp pro ile ser asp glu ile met asm gly thr ile 820 825 830 ala gly ile val ser gly ile ser ala ser glu ser leu leu ser gln 835 840 845 lys ser ata leu ile asp pro ala thr asm met val val gly glu phe 850 855 860 gly gly leu leu asn pro ala thr gly val met île pro gly phe leu 865 870 880 gly pro ser glu gln thr gln phe ser pro glu ile glu asp gly gly 885 890 895 ile ile pro pro glu val ala ala ala asn ala asp lys phe lys leu 900 905 910 ser ile pro pro ser val pro glu ser ile pro glu lys asp gln lys 915 920 925 ile asp ser ile ser glu leu met tyr asp ile glu ser gly arg leu 930 935 940 ite gly gin val ser tys arg pro ite pro gly ser ite ata gly asp 945 950 955 960 leu asn pro ite met tys thr pro thr gin thr asp ser val thr gly 965 970 985 lys pro ile asp pro thr thr gly leu pro phe asm pro pro thr gly 980 985 990 his leu ile asn pro thr asn asn asn thr met asp ser ser phe ala
995 1000 1005 gly ala tyr lys tyr ala val ser asn gly ile lys thr asp asn val 1010 1015 1020 tyr gly leu pro val gly glu ile thr gly leu pro lys asp pro gly 1025 1030 1035 1040 ser asp ile pro phe asn ser thr thr gly glu leu val asp pro ser 1045 1050 1055 thr gly lys pro ile asn asn ser thr ala gly ile val ser gly lys 1060 1065 1070 pro gly leu pro pro ile glu asp glu asn gly asn leu phe asp pro 1075 1080 1085 ser thr asn leu pro ile asp gly asn asn gln leu val asn pro glu 1090 1095 1100 thr asn ser thr val ser gly ser thr ser gly thr thr lys pro lys 1105 1110 1115 1120 pro gly ile pro val asn gly gly gly val val pro asp glu glu ala 1125 1130 1135 lys asp gln ala asp lys gly lys asp gly teu ile vat pro pro thr 1140 1145 1150 asn ser ile asn lys asp pro val thr asn thr gin tyr ser asn thr 1155 1160 1165 thr gly asn ile ile asn pro glu thr gly lys val ile pro gly ser 1170 1175 1180 leu pro gly ser leu asn tyr pro ser phe asn thr pro gln gln thr 1185 1190 1195 1200 asp glu ile thr gly lys pro val asp thr val thr gly leu pro tyr 1205 1210 1215 asp pro ser thr gly glu ile ile asp pro ala thr lys leu pro ile 1220 1225 1230 pro gly ser val ala gly asp glu île leu thr glu val leu asn île 1235 1240 1245 thr thr asp glu val thr gly leu pro ile asp leu glu thr gly leu 1250 1255 1260 pro arg asp pro val ser gly leu pro gln leu pro asn gly thr leu 1265 1270 1280 val asp pro ser asn lys lys pro ile pro gly ser his ser gly phe
1285 1290 1295 ile asn gly thr ser gly glu gln ser his glu lys asp pro ser thr 1300 1305 1310 gly lys pro leu asp pro asn thr gly leu his pro phe asp glu asp 1315 1320 1325 ser gly ser leu ile asn pro glu thr gly asp lys leu gln gly ser 1330 1335 1340 his ser gly thr phe met pro val pro gly lys pro gln gly glu asn 1345 1350 1355 1360 gly gly fle met thr pro glu gln ile leu glu ala leu asn lys leu 1365 1370 1375 pro thr ser asm glu val asm ile ser pro arg pro ser ser asp ala 1380 1385 1390 val pro asp arg pro thr asn thr trp trp asn lys ile ser gly gln 1395 1400 1405 thr tyr gln val asp gly lys lys thr ile leu gly ser ala ala ser 1410 1415 1420 val ile his thr ala leu gly thr pro thr gln thr asp pro thr thr 1425 1430 1435 1440 gly leu pro ser asp pro ser thr gly leu pro phe ile pro gly phe 1445 1450 1455 asn val leu val asp pro gln thr gly glu gln ile lys gly ser val 1460 1465 1470 pro tyr val ser leu tyr val lys glu lys asn ile val thr glu ala 1475 1480 1485 ala tyr gly leu pro val asp pro lys thr gly phe pro ile asp pro 1490 1495 1500 ile ser tyr leu pro phe ala lys asn gly glu leu ile asp pro ile
1505 1510 1515 1520
ser gly lys tyr phe ser gly ser ile ala gly phe ile ser gly lys
1525 1530 1535 ala gly ser gin ser lys ser ser asp glu ser gly asn pro ile asp 1540 1545 1550 pro ser thr asn met pro tyr asp pro lys gly gly lys leu ile asp 1555 1560 1565 pro glu ser gly ile ala ile asp asn ser val ser gly val phe ala 1570 1580 thr val pro gly thr ata ata pro lys lys gly gly val ite pro glu
1585 1590 1595 1600 ser val ala ala glu ala ala lys lys tyr phe ala ala asn val glu 1605 1610 1615 glu gly glu gly glu glu val pro pro pro glu ser ser ser 1620 1625 1630 ile ala ile gin ala ala gly gly ala ser ala ala val gly leu 1635 1640 1645 val ala ala val gly ala trp tyr ala ser arg asn arg gin glu gly 1650 1655 1660 glu asp asp asp tyr gin met asp leu lys gin asn met lys lys 1670 1675 1665 lys arg lys lys arg val met lys gln gin met lys leu leu leu gln 1685 1690 leu ser val ile his his ser gly thr ash leu lys arg arg lys asp 1705 1700 phe ser asn ser lys lys phe arg ile 1715

(7) INFORMATION FOR SEQ ID NO. 6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 503 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (vi) ORIGINAL SOURCE:
- (A) ORGANISM: Cryptosporidium parvum
- (ix) FEATURE
 - (A) NAME/KEY: antigen
 - (B) IDENTIFICATION METHOD: deduced
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO. 6:

ser lys gly gln leu phe asn pro val ser lys leu cys val arg leu 10 lys asp asn val val gly gly gly ala leu val leu asp asp cys arg
20 25 30 lys ala ser asp gly ser gly leu phe glu leu met pro asn asn gln
35 40 45 40 leu arg leu ala arg gly gly asn leu cys leu thr ser pro gly asp 50 55 60 tys pro gly val ala asn val ala leu asn ser ala ala ser ser thr 75 70 ser val val arg thr gly ile glu asn gly pro ala met ala val asp 85 90 95 gly lys asp thr ser tyr trp leu ser asp ser ser thr pro gly lys 100 105 110 asp ser ala asn val asn phe leu ser asp thr gly ser val thr lys 115 120 125 120 115 leu lys asp ile phe ile glu trp lys tyr pro ala ile asp phe asn
130 140 130 ile asp leu ser glu asn gly lys glu tyr gln thr gln val ser val 145 150 155 asn asn asn gly leu met ser thr thr tyr ser leu glu gly lys lys

ala arg tyr val lys ile gln met thr ile pro ser gln asp glu thr gly lys tyr val tyr gly ile lys gln val arg ile phe ser asn thr met arg ser thr val glu asp cys ser ser val lys gln his asn asp gly arg asp lys ile phe pro leu pro tyr asn gly asp asn phe ala pro gly leu leu lys ala his gly ile ser val lys asn arg leu asn glu leu gln glu leu ser gly lys val thr ser ile leu pro asn leu asp ala cys arg lys thr ser asp gly arg asp asn thr leu lys met gln ala thr lys leu gly phe leu ser glu lys leu glu lys leu thr ser asp tyr asn leu glu tyr lys phe thr lys pro ala leu gly gly ser glu leu tyr pro gly glu asp cys val ala ile lys asn asp lys thr gln glu ala ile ser gly phe tyr tyr val arg pro phe cys ser thr lys pro leu arg val tyr cys asp met asn thr gly asn thr ile tyr pro met glu met ser val his ser ser arg ala ala ser ser ala cys ala thr val gly leu lys pro leu leu leu arg asp lys lys glu ser val val gly ile lys lys met leu asn met met asn ile asn asp asn arg arg val ile pro leu thr his asp phe gly cys asp asn pro lys gly cys asn ser gln phe thr gln leu gly ser gly val glu glu phe val ala ala ser pro gln ala ala ala ser asn ser thr ser gly ala leu pro glu leu val leu cys ser thr asn thr asn leu lys his glu ser asn ala ile ser leu ser cys glu ser arg phe ser asp met lys val phe his leu asp

WHAT IS CLAIMED IS:

- 1. An anti-Cryptosporidium antibody binding to a Cryptosporidium peptide or glycopeptide antigen in a host.
 - 2. The antibody of claim 1 produced in vivo or in vitro.

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- 3. The antibody of claim 2, wherein the antibody is a monoclonal antibody, polyclonal antibody or chimeric construct.
- 10 4. The antibody of claim 3 capable of binding to a Cryptosporidium peptide antigen having Mr greater than 900.000.
- 5. The antibody of claim 3 capable of binding to a 15 Cryptosporidium peptide antigen having Mr between 50,000 and 100,000.
- 6. A method for treatment of Cryptosporidium infections comprising administering to a subject in need of such treatment an inhibitory amount of an anti-Cryptosporidium antibody binding to a Cryptosporidium peptide or glycopeptide antigen in a host.
- 7. The method of claim 6 wherein the antibody
 25 administered to the subject is produced in vivo or in vitro.
- 8. The method of claim 7, wherein the administered antibody is a monoclonal antibody, polyclonal antibody or chimeric construct.
- g. The method of claim 8 wherein the antibody administered to the subject is capable of binding to a cryptosporidium peptide antigen having Mr greater than 900,000.

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10. The method of claim 8 wherein the antibody administered to the subject is capable of binding to a Cryptosporidium peptide antigen having a Mr between 50,000 and 100,000.

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- 11. A method for active prophylaxis of Cryptosporidium infection comprising administering to a subject in need of such prophylaxis an amount of a Cryptosporidium peptide or glycopeptide antigen capable of binding to an anti-Cryptosporidium antibody, said amount of the antigen sufficient to elicit production of anti-Cryptosporidium antibodies.
- 12. The method of claim 11 wherein the antigen
 15 administered to the subject has an amino acid sequence SEQ
 1D NO: 5 or SEQ ID NO: 6.
- 13. The method of claim 12 wherein the Cryptosporidium parvum peptide antigen administered to the subject has an amino acid sequence SEQ ID NO: 5 and a Mr greater than 900,000.
- 14. The method of claim 12 wherein the Cryptosporidium parvum peptide antigen administered to the subject has a amino acid sequence SEQ ID NO: 6 and a Mr between 50,000 and 100,000.
- 15. A method for passive prophylaxis of Cryptosporidium infection comprising administering to a subject in need of such prophylaxis an amount of an anti-Cryptosporidium antibody sufficient to ameliorate the infection.
- 16. A vaccine for immunization of a human or animal host comprising an administration to the host a Cryptosporidium peptide or glycopeptide antigen, or DNA or RNA encoding said peptide or polypeptide antigen, capable of evoking production of anti-Cryptosporidium antibodies.

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- 17. The vaccine of claim 16 wherein the administration of the antigen, DNA or RNA evokes in the host an active immunity against Cryptosporidium infection.
- 18. The vaccine of claim 17 wherein in response to the vaccine administration the host produces cytokines, lymphokines, or antibodies, or acquires a cellular capacity to inhibit or retard Cryptosporidium infection in response to a contact with a Cryptosporidium antigen.

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- 19. The vaccine of claim 18 additionally containing an appropriate pharmaceutically acceptable adjuvant.
- 20. The vaccine of claim 19 producing an amount of anti15 Cryptosporidium antibodies sufficient to retard, inhibit or
 counter the infection in a host.
 - 21. The vaccine of claim 20 wherein the antigen has a sequence SEQ ID NO: 5 and a Mr greater than 900,000.

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- 22. The vaccine of claim 21 wherein the antigen has a sequence SEQ ID NO: 6 and a Mr between 50,000 and 100,000.
- 23. A method for passive immunization of a human or animal host comprising administration to the host of anti-Cryptosporidium antibodies produced against Cryptosporidium peptide or glycopeptide antigen, said antibodies capable of inhibiting or retarding Cryptosporidium infection.
- 30 24. The method of claim 23, wherein the antibody is monoclonal or polyclonal or a chimeric construct.
- 25. The method of claim 24 wherein the antibody is capable of binding to a Cryptosporidium peptide antigen 35 having Mr greater than 900 kDa.
 - 26. The method of claim 25 wherein the antibody is capable of binding to a Cryptosporidium peptide antigen having a Mr between 50,000 and 100,000.

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- 27. A Cryptosporidium antigen having an amino acid sequence SEQ ID NO: 5 or SEQ ID NO: 6, a mutant or variant thereof.
- 5 28. The antigen of claim 27 having the amino acid sequence SEQ ID NO: 5 additionally containing a N-terminal extension thereof.
- 29. The antigen of claim 22 capable of producing an 10 anti-Cryptosporidium antibody in vivo or in vitro.
 - 30. The antigen of claim 29 having an amino acid sequence SEQ ID NO: 6 additionally containing a N- terminal extension thereof.

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- 31. The antigen of claim 30 capable of producing an anti-Cryptosporidium antibody in vivo or in vitro.
- 32. A recombinant DNA or a fragment thereof having a sequence SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3, or SEQ ID NO: 4 or to a mutant or a variant thereof.
 - 33. The DNA of claim 32 additionally comprising a 5' extension.

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- 34. The DNA of claim 32 having a sequence SEQ ID NO: 1, a variant or mutant thereof.
- 35. The DNA of claim 34 additionally comprising a 5' extension.
 - 36. The DNA of claim 32 having a sequence SEQ ID NO: 2, a variant or mutant thereof.

35

37. The DNA of claim 36 additionally comprising a 5' extension.

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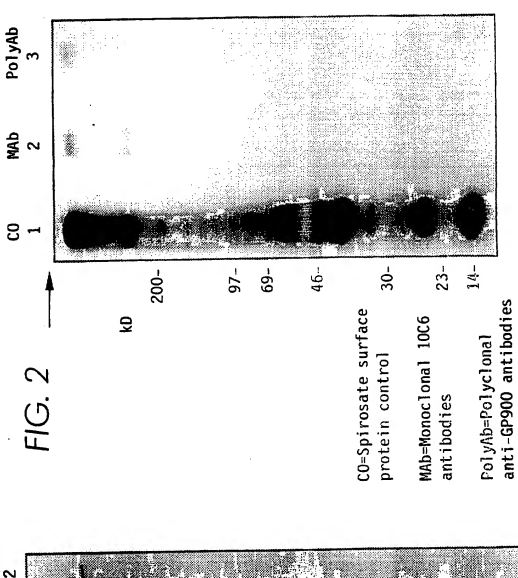
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- 38. The DNA of claim 32 having a sequence SEQ ID NO: 3, a variant or mutant thereof.
- 39. The DNA of claim 38 additionally comprising a 5' extension.
 - 40. The DNA of claim 32 having a sequence SEQ ID NO: 4. a mutant or variant thereof.
- 10 41. The DNA of claim 40 additionally comprising a 5' extension.
- 42. A recombinant RNA or a fragment thereof corresponding to a DNA sequence SEQ ID NO: 1, SEQ ID NO: 2, SEQ ID NO: 3 or SEQ ID NO: 4, or to a variant or mutant thereof.
 - 43. The RNA of claim 42 corresponding to a DNA sequence SEQ ID NO: 1, or a variant or mutant thereof.

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- 44. The RNA of claim 41 corresponding to a DNA sequence SEQ ID NO: 2, or a variant or mutant thereof.
- 45. The RNA of claim 41 corresponding to a DNA sequence 25 SEQ ID NO: 3, or a variant or mutant thereof.
 - 46. The RNA of claim 41 corresponding to a DNA sequence SEQ ID NO: 4, or a variant or mutant thereof.
- 30 47. A purified recombinant protein or fragment thereof having a sequence SEQ ID NO: 5 or SEQ ID NO: 6.

MAb PolyAb



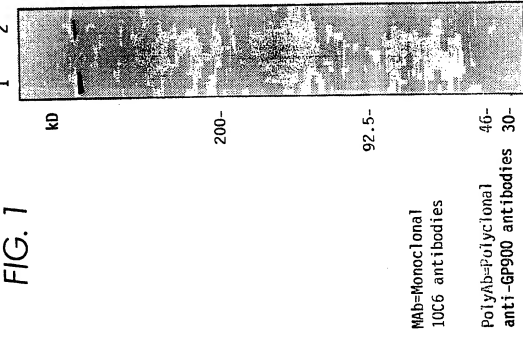
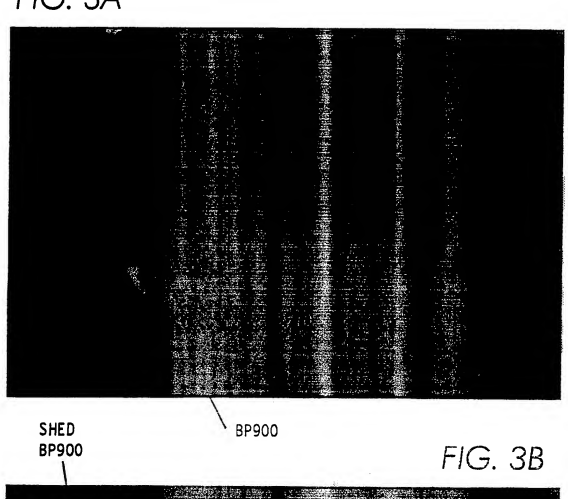
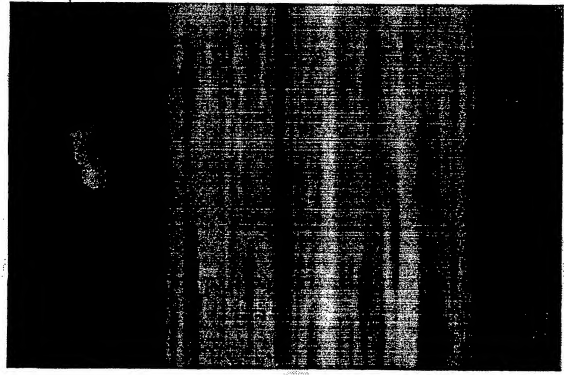
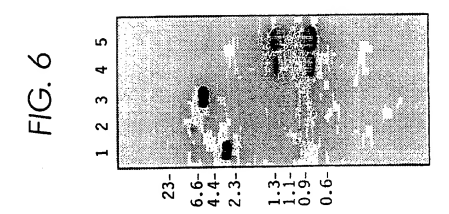


FIG. 3A







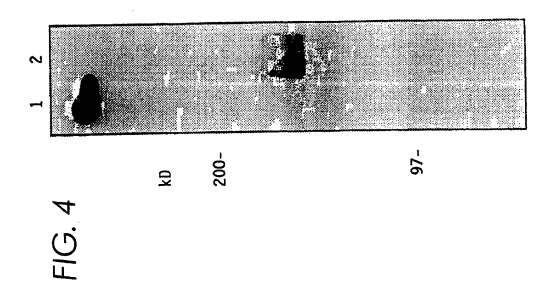
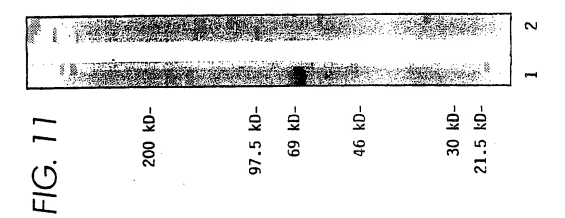


FIG. 5





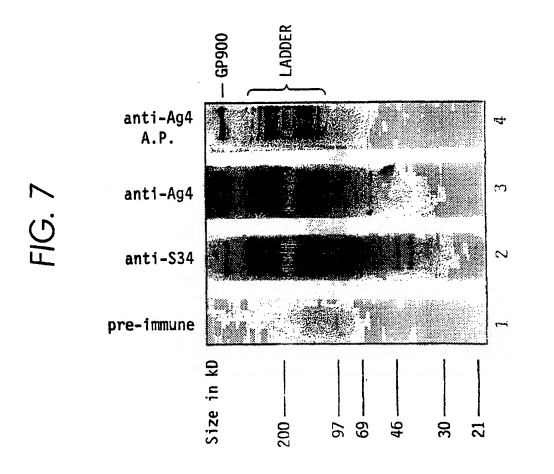
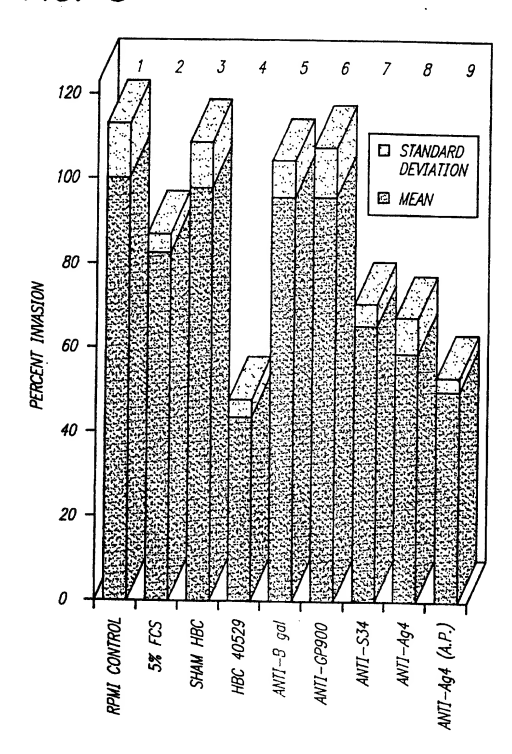
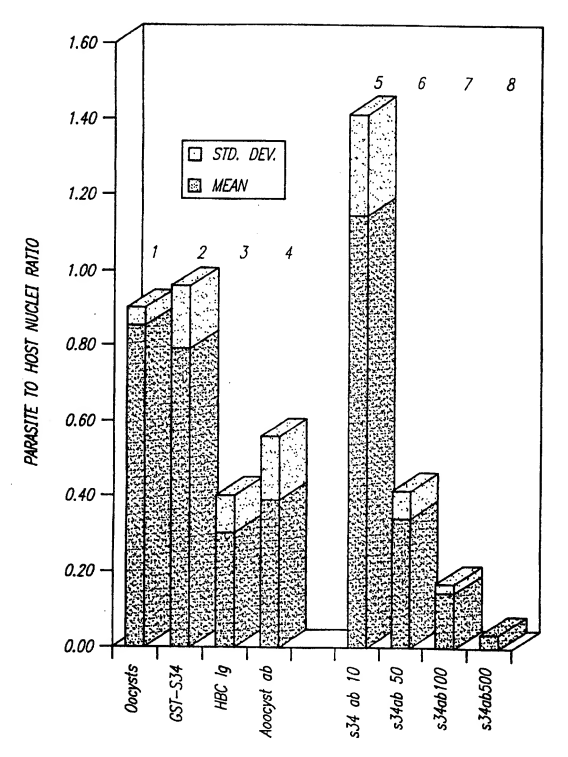


FIG. 8





ANTIBODIES AND CONTROLS

FIG. 9

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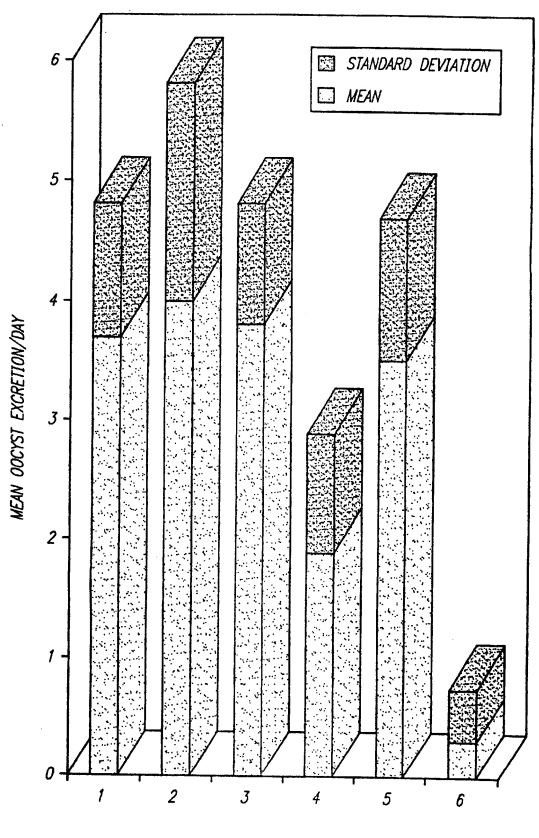


FIG. 10

FIG. 12

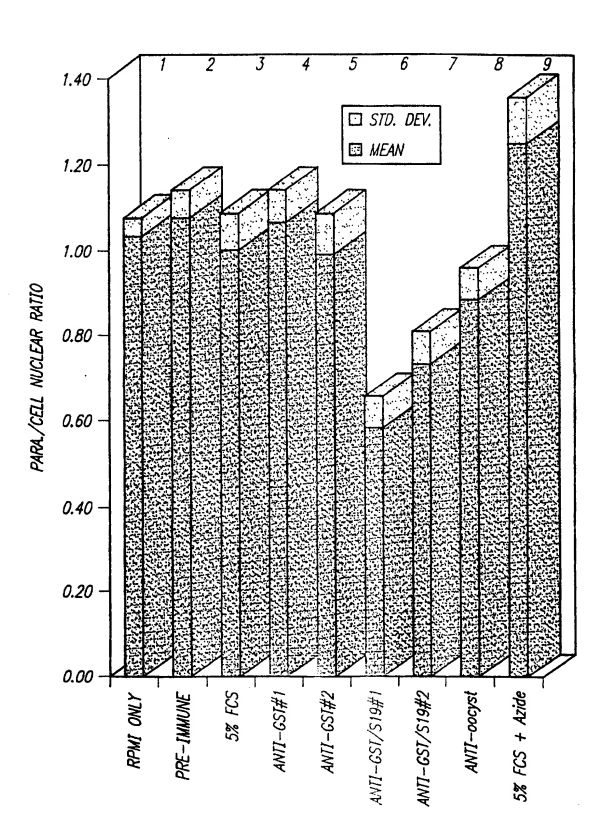
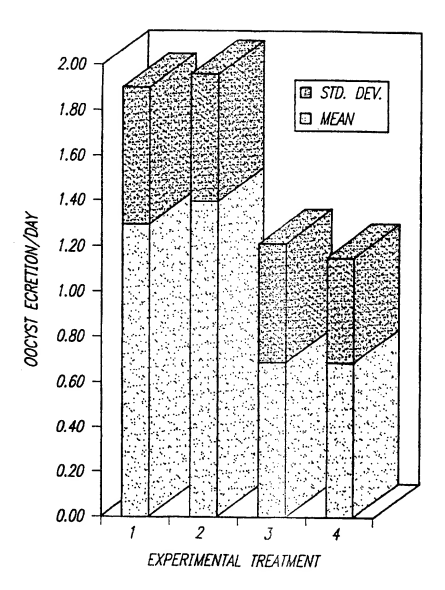


FIG. 13



INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/14104

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A61K 39/395, 39/40, 39/42, 39/44					
US CL :424/178.1; 514/44					
According to International Patent Classification (IPC) or to both national classification and IPC					
	LDS SEARCHED				
Minimum o	documentation searched (classification system follow	ed by classification symbols)			
U.S. : 424/178.1; 514/44					
Documenta	tion searched other than minimum documentation to the	ne extent that such documents are included	in the fields searched		
Electronic	data base consulted during the international search (name of data base and, where practicabl	c, search terms used)		
	ALOG, GENEBANK	•	,		
	rms: Cryptosporidium parvum, DNA, antibod?, vec	tor, protein?, glycoprotein?, treatment.	vaccin?. immunology?		
C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category*	Citation of document, with indication, where a		Relevant to claim No.		
Х, Р	US 5,643,772 A (PETERSEN et a document.	l) 01 July 1997, see whole	1-47		
X, P	US 5,591,434 A (JENKINS et al) (document.	1-3, 6-8			
X	RIGGS, M. W. et al. Neutralization-s on the surface of infectious Cryptosp The Journal of Immunology. 15 Aug pages 1340-1345, especially page 134	poridium parvum sporozoites. gust 1989, Vol. 143, No. 4,	1-3, 5		
X Further documents are listed in the continuation of Box C. See patent family annex.					
• Spe	soial entegories of cited documents:	"T" leter document published after the inte	mational filing date or priority		
	rument defining the general state of the art which is not considered be of particular relevance	date and not in conflict with the appli the principle or theory underlying the	cation but nited to understand		
	tier document published on or after the international filing date	"X" document of particular relevance: the	risimad investion comes to		
	pument which may throw doubts on priority claim(s) or which is	considered novel or cannot be consider when the document is taken alone	red to involve an inventive step		
cita	ed to establish the publication date of another citation or other scial reason (as specified)	"Y" document of perticular relevance; the			
	pument referring to an oral dischasure, use, exhibition or other	companies to madia an invantive	sten when the decomment is		
***		combined with one or more other such being obvious to a person skilled in the	documents, such combination		
"P" doc	nument published prior to the international filing date but later than priority date claimed	*&* document member of the same patent	family		
Date of the actual completion of the international search Date of mailing of the international search report					
29 SEPTEMBER 1997		1.4 NOV 1997			
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT		Authorized officer			
	ı, D.C. 20231	GINNY PORTNER	5/1		
Facsimile N		Telephone No. (703) 308-0196			

INTERNATIONAL SEARCH REPORT

International application No.
PCT/US97/14104

		PC17US97/141		
C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where appropriate, of the releva	int passages	Relevant to claim No.	
х Y	PEETERS, J.E. et al. Cryptosporidium parvum in calves: and immunoblot analysis of specific serum and local ant responses (Immunoglobulin A [IgA], IgG, and IgM) afte and experimental infections. Infection and Immunity. Jun Vol. 60, No. 6, pages 2309-2316, especially pages 2311-	ibody r natural ne 1992.	1-3, 5-8, 10, 15, 23-24, 26 	
 Ү	PETERSEN, C et al. Characterization of a >900,000 M, Cryptosporidium parvum sporozoite glycoprotein recogni. Protective hyperimmune bovine colostral immunoglobulii Infection and Immunity. December 1992, Vol. 60, No. 125132-5138, see whole document.	n.	1-4, 6-11, 15, 16- 20, 23-25, 29, 31 12, 13, 21, 27, 28, 29	
Y	PETERSEN, C. et al. Identification and Initial Characteri five Cryptosporidium parvum sporozoite antigen genes. It and Immunity. June 1992, Vol. 60, No. 6, pages 2343-23 especially pages 2345, table 1.	nfection	1-5, 16-22 27-31	
	Database EMBASE, No. 96007418, JENKINS, M. et al., and colostrum antibody responses induced by jet injection with DNA encoding a Cryptosporidium parvum antigen, Vaccine. 1995, Vol. 13, No. 17, pages 1658-1664, see enabstract.	of sheep abstract,	16-20	

INTERNATIONAL SEARCH REPORT

International application No. PCT/US97/14104

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s) 1-10, 15, and 23-26, drawn to antibodies and a method of using said antibodies.

Group II, claim(s) 11-14, 27-31, 47 and 16-22, drawn to antigen, method of immunizing, vaccines comprising said antigen.

Group III, claim(s) 32-46, and 16-22, drawn to DNA, RNA and compositions comprising said nucleic acids for the generation of an immune response.

The inventions listed as Groups I and II do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Group I is drawn to antibodies and methods of using said antibodies which evidence a different special technical feature from the antigen claimed in Group II; structurally the antibodies of claim 1 and the antigens of claim II differ in structure, and function and therefore differ in the special technical feature being claimed and therefore define different inventive concepts.

The inventions listed as Groups II and III do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Group II is drawn to antigens and their use in methods of treatment, Group III is drawn to DNA or RNA, and the use of said DNA or RNA in a method of treating a host using compositions comprizing the DNA or RNA; Groups II and III but differ in the three dimensional characteristics, molecular structure, mode of action and effect generated of the molecules of each group and therefore do not represent a single inventive concept.